## REMARKS

Claims 12, 14-20 and 31-34 are amended. Claim 35 is added. Claims 22-30 have been withdrawn from consideration by the Examiner. Claims 12-14, 16-21 and 31-35 are active and under consideration.

At the outset, Applicants wish to thank Examiner Marvich for the helpful and courteous discussion conducted with their U.S. representative, on March 17, 2009. The supporting remarks for patentability are consistent with the remarks made during the discussion with the Examiner.

Claims 12-15, 19-21 and 31-34 stand rejected under 35 USC 102(b) as being anticipated by *Poquet et al*, J. Bacteriology, 1998, Vol. 180, pages 1904-1912.

However, this reference fails to disclose or suggest the claimed invention.

Notably, <u>Poquet et al</u> (J. Bacteriology, 1998, Vol. 180: 1904-1912) disclose a plasmid pFUN comprising sequence accession number GenBank U95834 which contains the sequence named nlp3 (see attached copy of NCBI description of GenBank U95834). The sequence accession number U95834 consists of:

- i)  $p_{70}$  promoter;
- ii) a sequence encoding ZitR; and
- iii) part of sequence coding for ZitS.

This sequence has been cloned in the polylinker of pFUN as to obtain expression of a fusion peptide n1p3/ $\Delta$ Nuc. Therefore, <u>Poquet et al</u> disclose a plasmid containing the p<sub>Zn</sub> promoter (promoter sequence corresponding to SEQ ID NO: 1 of claimed expression cassette), a sequence encoding ZitR, the N-terminal part of ZitS fused with the sequence coding for  $\Delta$ Nuc, followed by the downstream restriction sites of the polylinker.

Amended claims 12 and 15 clearly relate to an expression cassette which does not contain

a sequence encoding any part of L. lactis **ZitS**. Consequently, the amended claims are clearly not anticipated by *Poquet et al.* 

It is also clear that <u>Poquet et al</u> would have failed to have rendered the claimed invention obvious at the time it was made. <u>Poquet et al</u> identified among others a sequence (U95834 GenBank sequence) which contains n1p3 coding sequence. <u>Poquet et al</u> are only interested in exported peptide n1p3, and are totally silent about a possible function of the other nucleotide sequences contained in U95834 GenBank sequence. Nothing in the disclosure of <u>Poquet et al</u> or in accession number U95834 report would have suggested to the artisan that any other sequences contained in U95834 sequence could have a regulating function or code for a protein. See the attached NCBI description of GenBank U95834.

Therefore the person skilled in the art would not have been motivated to isolate specifically the nucleotide sequence corresponding to  $p_{Zn}$  promoter and sequence encoding ZitR from sequence GenBank accession number U95834 in order to engineer an expression cassette which has the characteristic to regulate expression of a gene according to the concentration of zinc in the culture medium.

Further, <u>Poquet et al</u> would not have enabled one skilled in the art to do so in any event.

Hence, this ground of rejection is unsustainable and should be withdrawn.

Claims 12-21 and 31-34 stand rejected under 35 USC 112, first paragraph, as the present specification ostensibly does not provide enablement for any embodiment of expression cassettes other than that containing SEQ ID NO: 1 operably linked to nucleotides 357-794 of SEQ ID NO: 2 further operably linked to a restriction site.

However, it is believed that the present claims are fully enabled by the present specification for the following reasons. The comments set forth below refer to Annex 1 and Annex 2, copies of

which are attached to this response.

First, the definition "at least 80% identity with the *Lactococcus lactis* ZitR protein GENBANK AAK06214" does not encompass a broad variety of proteins.

Second, Annex 1 shows the results of a BLAST search performed against the nr database which includes all the known protein sequences of all living organisms. Among these sequences, the only ones which share at least 80% identity with ZitR of L. *lactis* subsp. Lactis II1403 (GenBank AAK06214) are ZitR proteins, namely the ZitR protein of L. *lactis* subsp. *cremoris* MG1363 (88% identity) and the ZitR protein of L. *lactis* subsp. *cremoris* SK11 (89% identity). As a matter of fact there is no protein having at least 80% identity with GenBank AAK06214 which is not a ZitR protein. The next proteins which have the higher homology with ZitR are Streptococcus proteins which have at most 54% identity with GenBank AAK06214. Therefore, it is very unlikely that one can isolate a protein having 80% identity or more with GenBank AAK06214 which is not a ZitR protein.

Annex 2 shows the results of a BLAST search performed against the whole genome sequences of L. *lactis* subsp. Lactis II1403, L. *lactis* subsp. *cremoris* MG1363 and L. *lactis* subsp. *cremoris* SK11. There is no lactococcal protein, other than the ZitR proteins, having more than 38% identity with GenBank AAK06214.

Therefore using a probe derived from the sequence encoding GenBank AAK06214, or from nucleotides 357 to 794 of SEQ ID NO:9 of the instant application for screening a lactococcal DNA library, one of skill in the art would be able to easily discriminate a sequence encoding a ZitR protein from other lactococcal sequences.

Moreover, only the whole genome sequence of L. *lactis* subsp. Lactis II1403 (which was published in 2001) was available before the invention was made. The genome sequences of *Lactococcus lactis* subsp. *cremoris* SK11 and *Lactococcus lactis* subsp. *cremoris* MG1363 were only available in GenBank in 2006 and 2007 respectively. It is pointed out, however, that besides indicating GenBank AAK06214 as the reference sequence for a ZitR protein, the present application discloses another sequence encoding a ZitR protein, namely nucleotides 357 to 794 of SEQ ID NO:9, which encodes the ZitR protein of *lactis* subsp. *cremoris* MG1363.

Actually, the knowledge of the lactococcal genome sequences is not necessary to practice the claimed invention. As indicated above, one of ordinary skill in the art could have easily obtained polynucleotides encoding lactococcal ZitR proteins by screening a DNA library of a *Lactococcus* with a probe derived from the sequence encoding GenBank AAK06214, or from nucleotides 357 to 794 of SEQ ID NO:9.

BLAST results are provided herewith solely to evidence that the lactococcal genome does not contain any protein having more than 80% identity with GenBank AAK06214 which is not a ZitR protein.

Second, the two BLAST searches and results thereof noted above indicate that the present specification does in fact satisfy both the statutory enablement requirement under 35 USC 112, first paragraph, and the case law tests therefor cited by the Examiner.

Specifically, under the case law tests for enablement articulated in <u>In re Wright</u> and <u>In re Fisher</u>, cited by the Examiner, it is clear that since nucleotide sequences having more than 80% identity with GENBANK AAK06214 could easily be obtained by routine screening with a probe (as described above), undue experimentation would not be required.

Claims 12-16, 18-20 and 31-34 are objected to.

However, in view of the above claim amendments, this ground of rejection is deemed moot.

The specification stands objected to.

However, attached to this response is a Substitute Specification which meets the requirements of 37 CFR 1.77(b).

Finally, Applicants hereby respond to the specific assertions set forth in the Notice of Non-Compliance of July 7, 2009.

First, claims 16, 18 and 20 have been amended as suggested by the Examiner. Claim 13 has been corrected.

Second, it is urged respectfully that claim 15 is <u>not</u> a duplicate of Claim 12. Notably claim 15 does not require element b) of claim 12, i.e., a sequence encoding a polypeptide with at least 80% identity with the *Lactococcus lactis* Zit R protein, placed under the transcriptional control of the promoter, and wherein the polypeptide is obtained from Lactococcus.

Third, Applicants have, in fact, addresses the rejection of claims 12-21 and 31-34 under 35 USC 112, first paragraph. See preceding pages 10-12 of this Amendment, wherein it is explained why these claims are enabled by the present specification.

## **CONCLUSION**

Accordingly, in view of all of the above, it is believed that this application is now in condition for allowance. Early notice to this effect is earnestly solicited.

Applicant hereby petitions for the Commissioner to charge any additional fees or any underpayment of fees which may be required to maintain the pendency of this case or credit any overpayment to Deposit Account No. 14-0112.

Respectfully submitted, THE NATH LAW GROUP

William E. Beaumont Registration No. 30,996 Customer No. 20529

Date: April 7, 2009

THE NATH LAW GROUP

112 S. West Street

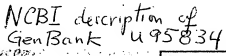
Alexandria, Virginia 22314

Tel: (703) 548-6284 Fax: (703) 683-8396

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All Databases

PubMed

Protein

Genome

Taxonomy

Search Nucleotide

for

Go

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Limits

Preview/Index

History

Format: GenBank FASTA Graphics More Formats ▼

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Links V

GenBank: U95834.1

Lactococcus lactis putative lipoprotein Nlp3 precursor, gene, partial cds

Features Sequence

LOCUS

LLU95834

945 bp

DNA

BCT 24-APR-1998 linear

DEFINITION Lactococcus lactis putative lipoprotein Nlp3

precursor, gene,

partial cds.

ACCESSION

U95834

VERSION

U95834.1 GI:3043865

**KEYWORDS** 

SOURCE

Lactococcus lactis subsp. cremoris MG1363

ORGANISM Lactococcus lactis subsp. cremoris MG1363

Bacteria; Firmicutes; Lactobacillales;

Streptococcaceae;

Lactococcus.

REFERENCE 1 (bases 1 to 945)

AUTHORS Poquet, I., Ehrlich, S.D. and Gruss, A.

TITLE An export-specific reporter designed for gram-

positive bacteria:

application to Lactococcus lactis

JOURNAL J. Bacteriol. 180 (7), 1904-1912 (1998)

PUBMED 9537391

REFERENCE (bases 1 to 945)

AUTHORS Poquet, I. and Gruss, A.

TITLE Direct Submission

JOURNAL Submitted (31-MAR-1997) Laboratoire de Genetique

Appliquee-URLGA,

Institut National de la Recherche Agronomique,

CRJ, Jouy en Josas

78352, France

**FEATURES** 

Location/Qualifiers

source

/organism="Lactococcus lactis subsp.

cremoris MG1363"

/mol\_type="genomic DNA"

/strain="MG1363"

/db xref="taxon:416870"

complement (<1..330)

/note="identified as a fusion to a

signal peptide-less

form of the staphylococcal nuclease

reporter which

CDS

displays nuclease activity; similar to

S. pneumoniae

adhesion protein, Swiss-Prot Accession

Number P42363"

Change Region Shown

**Customize View** 

Pick Primers

Design and test primers for this sequence using Primer-BLAST.

**Recent Activity** 

Turn Off Clear :

Lactococcus lactis putative lipoprotein

<u>U95834</u> (1) Mucleolide

All links from this record

Full text in PMC

Protein

PubMed

Taxonomy

Related Sequences

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peptide"
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actcctttat tattcacttt
       61 cagageceat ttttcaaggt catcagaatt ataaataata
gcttgagcat ttgtcattgt
      121 tgcaacttgc ttggcactcg gttcaaattc gtgaacttct
tgattcgccg gaacaatatt
      181 ttcaatttta accttatctc caacaatcgc tttcgtaaat
tcatacatcg gctcaaaagt
      241 tgtcacaact tctggtttgt ctgctgtttt ttgacaacca
gcaagaagta aaactgccgg
      301 aatagcaaat aacatcaata ttttcttcat cgaaactcct
ccgtaagtac tgataagaat
      361 tgacttatca ctttttgttc ttcgtcagta aatttgtctc
ctaattcttg gtaggtactt
      421 agagttttct catgatgagc agcatgttct ttagcaactg
gaattgcttt ttctgtcagg
      481 ctccaaagga ctacgcgttc gtcatttgtt gcccgacttg
atttaatcag ttcttgctct
      541 tgtaattttt tgagagettt agttaceget getggegaaa
tcttgagttg ctcggcaatt
      601 ctcgcgtttg tcgaaacctc tgcagctaga atcattaaga
tatgttcttg cgtgcttgtt
      661 agcttaacat tactttngca ttcgccgagt aatatttcat
gcttgttttn tgcaaactgc
      721 ataattgccc caagaaactg gtcgatttga tttgctaaan
tcatagtatt ttgtcctcca
      781 taattagttt actggttaat tatatagcaa agtaaaaata
atgtcaacca gttaacatta
      841 ttttttactt ttttatttga aaaatccttc catccttaag
ccgaacatca aaacaataag
      901 ttttattact aacagtttga ccaccgctag ttgaaccagc tgaca
```

11

Write to the Help Desk
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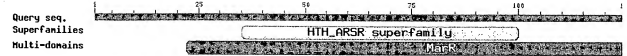
Length adjustment 109 Effective length of query 36

Effective length of database 1873821432 Effective search space 67457571552 Effective search space used 67457571552

Graphic Summary

Show Conserved Domains

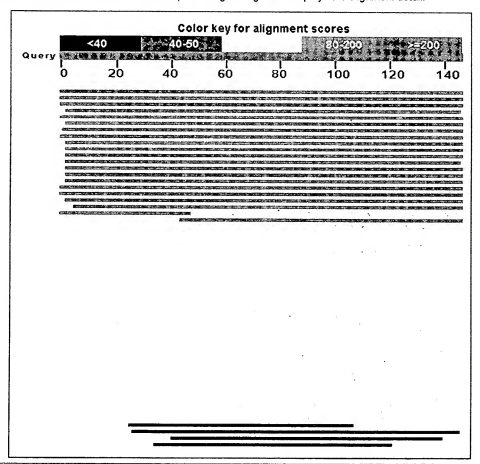
Putative conserved domains have been detected, click on the image below for detailed results.



Distribution of 100 Blast Hits on the Query Sequence

[?]

An overview of the database sequences aligned to the query sequence is shown. The score of each alignment is indicated by one of five different colors, which divides the range of scores into five groups. Multiple alignments on the same database sequence are connected by a striped line. Mousing over a hit sequence causes the definition and score to be shown in the window at the top, clicking on a hit sequence takes the user to the associated alignments. New: This graphic is an overview of database sequences aligned to the query sequence. Alignments are color-coded by score, within one of five score ranges. Multiple alignments on the same database sequence are connected by a dashed line. Mousing over an alignment shows the alignment definition and score in the box at the top. Clicking an alignment displays the alignment detail.



## Descriptions

Sequences producing significant alignments:	Score (Bits)	E Value
	,,	
<pre>ref[NF 168273.1] zinc transport transcription regulator [Lact</pre>	7501	1e-77 €
ref % \$11979 transcriptional regulator [Lactococcus lacti	258	1e-70 <b>G</b>
ref[[YF 001033643.1] transcriptional regulator of the zit oper	267	2e-70 <b>G</b>
<pre>ref[YF 001034147.1] multiple antibiotic resistance operon tra ref[NF 702293.1] putative transcriptional regulator [Streptoc</pre>	154	2e-36 <b>G</b> 1e-33 <b>G</b>
<u>401 YP 002122433.1</u> transcriptional repressor AdcR for Zn(2+)	145 143	3e-33 <b>G</b>
Lef AP 606407.11 putative repressor protein [Streptococcus py	191	1e-32 <b>G</b>
ref. ZP_02920004.: hypothetical protein STRINF_00865 [Strepto	1.4	2e-32
<pre>ref.NP 358489.1</pre> putative repressor protein [Streptococcus py	_14	2e-32
ref YP 002551480.1 Mark-family regulatory protein [Streptoco	138	1e-31 G
<pre>ref.NP 687190.11 adc operon repressor AdcR [Streptococcus aga ref.YF 001401005.1 repressor protein adcR [Streptococcus gor</pre>	<u> 136</u> 132	5e-31 <b>G</b> 4e-30 <b>G</b>
repressor protein adcR [Streptococcus gor ref 7F 01817760.1] adc operon repressor AdcR [Streptococcus p	132	4e-30 🚨 6e-30
ref.RF 350066.   adc operon repressor AdcR (Streptococcus pne	13:	2e-29 😉
$\underline{\text{ref-NI-346556.1}_1}  \text{adc operon repressor AdcR [Streptococcus pne}$	130	2e-29 <b>G</b>
<pre>ref.SP_138785.3* zinc transport transcriptional repressor [St</pre>	129	5e-29 <b>G</b>
ref WE 819746.11 zinc transport transcriptional repressor [St	128	1e-28 <b>G</b>
<pre>transcriptional regulator [Streptococcus tef 2P 03625066.1 transcriptional regulator, MarR family [St</pre>	<u>110</u>	7e-25 <b>G</b> 4e-23
emb_CAATE212.1 hypothetical protein [Lactococcus lactis subs	2 4	1e-17
COG1846: Transcriptional regulators [Strep	24.3	3e-17 5e-10 <b>G</b>
<u>ref:YF 001485459.1</u> Mark family transcriptional regulator [Ba ref:YF 001199572.1] transcriptional regulator [Streptococcus	67.0	5e-10 G 6e-09 G
ref YF 187315.11 Mark family transcriptional regulator [Staph	63.2 63.2	7e-09 <b>G</b>
reflyF 001777305.11 transcriptional repressor for Zn(2+)-resp	52.8	8e-09 <b>G</b>
raf UF 501177.1 hypothetical protein SAOUHSC 02819 [Staphylo	52.2	8e-09 <b>C</b>
ref:SF 03563431.1 Mark family transcriptional regulator [Sta	51.2	3e-08
<u>reflyP 1/4450.11</u> MarR family transcriptional regulator [Bacil	<u>6C.5</u>	4e-08 G
<u>refixP 080661.1</u> transcriptional regulator YvnA [Bacillus lic refixP 254603.1] hypothetical protein pSHaeC05 [Staphylococcu	<u>59.3</u>	1e-07 <b>G</b> 1e-07 <b>G</b>
refixp 093087.1 YvnA [Bacillus licheniformis ATCC 14580] >gb	98.3 59.3	1e-07 G
refize 00738634.1  Transcriptional regulator, Mark family [Ba	55.8	9e-07
ref. ZF 02613091.1 transcriptional regulator, MarR family [Cl	<u>54.3</u>	3e-06
ref/YE 001153547.1: Mark family transcriptional regulator [Cl ref/YE 001760642.1: Mark family transcriptional regulator [Cl	54.3	3e-06 <b>G</b> 3e-06 <b>G</b>
refixE 001053131.1] Mark family transcriptional regulator [La	54 <u>.3</u> 53.0	4e-06 G
ref[3F 02617530.1] transcriptional regulator, MarR family [Cl	53.9 53.9	4e-06
$\underline{\text{ref}_{1}\text{YP-861960.1}}  \text{MarR family transcriptional regulator [Grame}$	53,9	4e-06 G
refine 391388.11 hypothetical protein BSU35080 [Bacillus subt Lef] 2P 02994615.11 hypothetical protein CLOSPO 01734 [Clostri	53.5 53.5	5e-06 <b>G</b> 6e-06
refize 01994138.1; hypothetical protein DORLON_00120 [Dorea 1	53.1	6e-06
<pre>ref[ZP 03294054.1] hypothetical protein CLOHIR_02015 [Clostri ref[ZP_02011336.1] hypothetical protein CLOBAR_00939 [Clostri</pre>	53.1	7e-06 1e-05
ref:YP_001390364.1  MarR family transcriptional regulator [Cl	52.4 52.4	1e-05 G
ref: YF 001786497. [ Mark family transcriptional regulator [Cl	51.5	2e-05 <b>G</b>
$\underline{r}$ -f $  T     S08775.1  $ transcriptional regulator (Lactococcus lacti	31.2	2e-05 <b>G</b>
$\underline{\text{ref:YP 001898248.3}}  \text{MarR family transcriptional regulator [Ly}$	51.2	3e-05 <b>€</b>
<pre>ref(NP 266864.1) transcription regulator [Lactococcus lactis ref(ZP 01723095.1) transcriptional regulator, Mark family pro</pre>	51.2	3e-05 G
restrict and 17.11 transcriptional regulator (Lactococcus lacti	<u>50.4</u> 50.4	4e-05 5e-05 <b>G</b>
refine 786298.11 transcription regulator [Lactobacillus plant	50.1	6e-05 G
refiyP 757477.1] MarR family transcriptional regulator [Maric	50.1	6e-05 <b>€</b>
$\underline{\mathtt{ref}(\mathtt{NP} 391385,1)}  hypothetical \ \ protein \ \ BSU35050 \ \ [Bacillus subt$	50.1	6e-05 G
$\underline{\mathtt{ref}:\mathtt{NP}\ 111581.1!}  \mathtt{transcription}\ \mathtt{regulator}\ \mathtt{(SlyA-related)}\ \mathtt{[Ther}$	49.7	7e-05 <b>G</b>
ref:NF 419215.1; Mark family transcriptional regulator [Caulo	43.3	9e-05 G
ref YF 001031482.1; MarR family transcriptional regulator [La	13.3	1e-04 G 1e-04 G
ref NP 603578.1. Mark family transcriptional regulator [Fusob ref NP 138915.1. Mark family transcriptional regulator [Strep	<u> 1</u> 8.9 48.9	1e-04 G
ref. (%_000165566.); possible Mark family transcriptional regulator (strep	30.9 48.9	1e-04 G
Transcriptional regulator, Mark family [Fu	46.9	1e-04

# **BLAST Basic Local Alignment Search Tool**

Edit and Resubmit Save Search Strategies Formatting options Download

### gb|AAK06214| (145 letters)

Results for: | gb|AAK06214.1 zinc transport transcriptional regulator [Lactococcus lactis subsp. lactis II1403](145aa)

Your BLAST job specified more than one input sequence. This box lets you choose which input sequence to show BLAST results for.

#### Query ID

gi|12725171|gb|AAK06214.1|AE006439\_11

#### Description

zinc transport transcriptional regulator [Lactococcus lactis subsp. lactis II1403]

### Molecule type

amino acid

### **Query Length**

145

#### **Database Name**

nr

#### Description

All non-redundant GenBank CDS translations+PDB+SwissProt+PIR+PRF excluding environmental samples from WGS projects

### Program

BLASTP 2.2.19+ Citation

#### Reference

Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402.

Reference - compositional score matrix adjustment

Stephen F. Altschul, John C. Wootton, E. Michael Gertz, Richa Agarwala, Aleksandr Morgulis, Alejandro A. Schäffer, and Yi-Kuo Yu (2005) "Protein database searches using compositionally adjusted substitution matrices", FEBS J. 272:5101-5109. Other reports: Search Summary [Taxonomy reports] [Distance tree of results]

## **Search Parameters**

Program	blastp
Word size	3
Expect value	10
Hitlist size	100
Gapcosts	11,1
Matrix	BLOSUM62
Threshold	11
Composition-based stats	2
Filter string	F
Genetic Code	1
Window Size	40

#### **Database**

Posted date Mar 2, 2009 5:57 PM
Number of letters 2,739,991,458
Number of sequences 7,946,514
Entrez query none

## Karlin-Altschul statistics

Params	Ungapped	Gapped
Lambda	0.310848	0.267
K	0.124161	0.041
Н	0.32399	0.14

#### **Results Statistics**

```
Annex 1
                                                                                     1e-04 G
ref YP 819312.11 Mark family transcriptional regulator [Strep... 46.6
                                                                                     2e-04 G
ter. YP 079349.1| transcriptional regulator YvmB [Bacillus lic...
ref 2P 01219563.11 putative transcriptional regulator, MarR f...
                                                                                     2e-04
                                                                                     2e-04 G
ref YP 091766.1! YvmB [Bacillus licheniformis ATCC 14580] >gb... 45.1
                                                                                     2e-04 G
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201 ZP 00990845.1 hypothetical transcriptional regulator, Ma... 48.1
                                                                                     2e-04
                                                                                     3e-04 G
gol.YF 001316891.1; Mark family transcriptional regulator [Al... 47.8
                                                                                     3e-04 G
<u> 191 YP 001836303.11 MarR family transcriptional regulator [Le... 47.8</u>
                                                                                     4e-04 G
ref 7E 108511.1) putative transcriptional regulatory protein ... 47.4
                                                                                     4e-04 €
ref [F_02)961889.1: Transcriptional regulator, mark family [L... 47.4
\frac{\text{ref.} \ \text{YF.} \ 001310774.1)}{\text{ref.} \ \text{ZP.} \ 07138888.1)} \ \ \text{MarR family transcriptional regulator} \ \ \frac{47.4}{47.6} hypothetical transcriptional regulator, Ma... \frac{47.4}{47.6}
                                                                                     4e-04 G
                                                                                     5e-04
                                                                                     5e-04 G
LOT-YP 140799. 1 Mark family transcriptional regulator [Strep... 4...(
                                                                                     6e-04 C
<u>ref:rP 143939.11</u> MarR family transcriptional regulator [Therm... 46.6
                                                                                     7e-04 G
16f(xP 299371.1) regulatory protein, MarR [Ralstonia eutropha... 98.6
                                                                                     7e-04 G
hefrip 001707422.1 Mark family transcriptional regulator [Cl... 56.5
Tet ZP 02950515.1 transcriptional regulator, MarR family [Cl...
                                                                            48.3
                                                                                     8e-04
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                                                                            46.2
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                                                                            10.2
                                                                                     0.001
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ref CF 02.31766.1 transcriptional regulator, MarR family pro...
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col CP 602386 3.1, transcriptional regulator, Mark family [Ba...
                                                                                     0.001
                                                                           40.5
                                                                                     0.001
transcriptional regulator, MarR family [C...
                                                                            15.5
166 ZP 075-7491.1
                      hypothetical protein BLAHAN_01259 [Blautia...
                                                                                     0.001
ref ZP 02891041.1 transcriptional regulator, Mark family [Bu...
                                                                            45.4
                                                                                     0.001
                                                                                     0.001 G
ref:YF 001808132.1| MarR family transcriptional regulator [Bu...
ref:ZF 003006301.1| transcriptional regulator, MarR family [Bu...
ref:ZF 00360596.1]
                                                                            45.4
                                                                            45.4
                                                                                     0.001
                                                                            45.4
                                                                                     0.001
                                                                                     0.002 G
ref[NP 377055.1] Mark family transcriptional regulator [Bacil...
                                                                            45.4
                                                                                     0.002 G
ref[NP 780475.1] Mark family transcriptional regulator [Clost...
                                                                            45.4
ref(3P 03570809.1; transcriptional regulator, MarR family [Bu...
                                                                                     0.002
                                                                            45.I
                                                                                     0.002 C
ref[xP 001119315.1] MarR family transcriptional regulator [Bu... 45.1
                                                                                     0.002 G
\frac{\texttt{tef}(\texttt{xP}) \cdot \texttt{002418276.1}}{\texttt{ref}(\texttt{ZP}) \cdot \texttt{02355848.1}} \quad \text{Histone acetyltransferase HPA2 and relate...} \quad \frac{45.1}{45.1} \text{transcriptional regulator, MarR family pro...} \quad \frac{45.1}{45.1}
                                                                                     0.002
                                                                                     0.002 G
RefigP 443075.1 Mark family transcriptional regulator [Burkh...
                                                                            44,7
refixE 02633998.1 transcriptional regulator, MarR family [Cl... 44.3
                                                                                     0.002
                                                                                     0.002 G
ref YF 001374721.1: MarR family transcriptional regulator [Ba... 44.7
                                                                                     0.002 😉
ref[[Mr. 848464.1] transcriptional regulator [Leuconostoc mesen... 44.7
ref[SP 00027443.1] hypothetical protein EUBVEN_02713 [Eubacte... 44.7
                                                                                     0.003
```

Alignments Select All Get selected sequences Distance tree of results

LGNKFTDEEQEVISKFLSALTEEFQ 145

```
lactis Tl1403]
 gb|AAK06214.1|AE006439 11 C zinc transport transcriptional regulator [Lactococcus lactis
subsp. lactis Il1403]
Length=145
GENE ID: 1115793 zitR | zinc transport transcription regulator [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 291 bits (744), Expect = 1e-77, Method: Compositional matrix adjust. Identities = 145/145 (100%), Positives = 145/145 (100%), Gaps = 0/145 (0%)
             MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL
MSLANOIDOFLGTIMOFAENKHEILLGKCESDVKLTSTOEHILMLLAEOISTNAKIAEKL
Query 1
Sbjct 1
             MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL
             KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
Query 61
                                                                                     120
             KISPAAVTKALKKLÕEÕELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYÕE
             KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
Sbjct 61
             LGNKFTDEEQEVISKFLSALTEEFQ 145
Query 121
             LGNKFTDEEÕEVISKFLSALTEEFÖ
```

```
GENE ID: 4433026 LACR 2420 | transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 268 bits (685), Expect = 1e-70, Method: Compositional matrix adjust. Identities = 130/145 (89%), Positives = 141/145 (97%), Gaps = 0/145 (0%)
                  MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL
Querv 1
                 MSLANQIDOFLG IMOFAENKHEILLG+CES+VKLTSTOEHILM+LA ++STNA+IAE+L
MSLANQIDOFLGAIMOFAENKHEILLGECESNVKLTSTOEHILMILAAEVSTNARIAEQL
Sbict 1
                                                                                                                60
                  KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
                                                                                                               120
Query 61
                 KISPAAVTKALKKLÕEÕELIKSSRATNDERVVLWSLTEKAVPVAKEHA HHEKTLSTYÕE
KISPAAVTKALKKLÕEÕELIKSSRATNDERVVLWSLTEKAVPVAKEHAAHHEKTLSTYÕE
Sbjct 61
                                                                                                               120
                 LGNKFTDEEQEVISKFLSALTEEFQ 145
LG+KFTDEEQ+VIS+FLS LTEEF+
LGDKFTDEEQKVISQFLSVLTEEFR 145
Query 121
Sbjct 121
>ref|YP 001033643.1| G transcriptional regulator of the zit operon (Lactococcus lactis
subsp. Cremoris MG1363]
 emb|CAL98965.1|  transcriptional regulator of the zit operon [Lactococcus lactis
subsp. cremoris MG1363]
Length=145
GENE ID: 4799067 zitR | transcriptional regulator of the zit operon [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 267 bits (682), Expect = 2e-70, Method: Compositional matrix adjust. Identities = 129/145 (88%), Positives = 141/145 (97%), Gaps = 0/145 (0%)
                  MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL
Query 1
                 MSLANQIDOFLG IMOFAENKHEILLG+CES+VKLTSTOEHILM+LA ++STNA+IAE+L
MSLANQIDOFLGAIMOFAENKHEILLGECESNVKLTSTOEHILMILAAEVSTNARIAEQL
Sbjct 1
                                                                                                                60
                 KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA+PVAKEHA HHEKTLSTYQE
KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAIPVAKEHAAHHEKTLSTYQE
Ouerv 61
Sbjct 61
                 LGNKFTDEEQEVISKFLSALTEEFQ 145
LG+KFTDEEQ+VIS+FLS LTEEF+
LGDKFTDEEQKVISQFLSVLTEEFR 145
Query 121
Sbjct 121
>ref|YP_001034147.1|  multiple antibiotic resistance operon transcriptional repressor (MarR), putative [Streptococcus sanguinis SK36]
gb|ABN43597.1|  Multiple antibiotic resistance operon transcriptional repressor
(Mark), putative [Streptococcus sanguinis SK36]
Length=147
 GENE ID: 4806188 adcR | multiple antibiotic resistance operon transcriptional
repressor (MarR). putative [Streptococcus sanguinis SK36] (10 or fewer PubMed links)
 Score = 154 bits (389), Expect = 2e-36, Method: Compositional matrix adjust. Identities = 78/142 (54%), Positives = 102/142 (71%), Gaps = 0/142 (0%)
                 LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKI
LA +IDQFL ++ AEN+HEIL+G C SDV LT+TQEHILMLL+E+ TN+ +A+KL +
LAQKIDQFLNEVILKAENQHEILIGSCTSDVPLTNTQEHILMLLSEESLTNSDLAKKLNV
Query 3
Sbict 4
Query 63
                  {\tt SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG}
                 S AAVTKA+K L QE++++ + D RV + LTE A P+AKEH HH TL TYQ+L SQAAVTKAVKSLARQEMLQAFKDKRDARVTFYRLTELAQPIAKEHOHHHAHTLETYQKLA
Sbjct 64
Query 123
                 NKFTDEEQEVISKFLSALTEEF 144
                 +F+ EQ VI+KFL AL E
EQFSASEQAVIAKFLEALVGEI 145
Sbict 124
>ref|NP_722293.1|  putative transcriptional regulator [Streptococcus mutans UA159]
 gb|AAN59599.1|AE015022 6 G putative transcriptional regulator [Streptococcus mutans UA159]
Length=148
GENE ID: 1029245 SMU.1995c | putative transcriptional regulator
[Streptococcus mutans UA159] (10 or fewer PubMed links)
 Score = 145 bits (365), Expect = 1e-33, Method: Compositional matrix adjust. Identities = 76/146 (52%), Positives = 105/146 (71%), Gaps = 1/146 (0%)
                 Ouerv 1
Sbjct 1
                 Query
Sbjct 61
                 ELGNKFTDEEQEVISKFLSALTEEFQ 145
+ KF+ +E+ VIS+FL+ALT+E +
RISQKFSQKEKSVISRFLTALTKELE 146
Query 120
Sbict 121
```

Anney '

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>ref|YP 002122498.1|  transcriptional repressor AdcR for Zn(2+)-responsive expression [Streptococcus equi subsp. zooepidemicus MGCS10565]

gb|ACG61485.1|  transcriptional repressor AdcR for Zn(2+)-responsive expression [Streptococcus equi subsp. zooepidemicus MGCS10565]
Length=147
GENE ID: 6760627 adcR | transcriptional repressor AdcR for Zn(2+)-responsive expression [Streptococcus equi subsp. zooepidemicus MGCS10565]
 Score = 143 bits (361), Expect = 3e-33, Method: Compositional matrix adjust. Identities = 70/143 (48%), Positives = 102/143 (71%), Gaps = 0/143 (0%)
               LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKI
Query 3
              L ++D + I+ AEN+HE+L G C+SDVKLT+TQEHILMLL+++ TN +A++L I LEKKLDNLVNRILLKAENQHELLFGACQSDVKLTNTQEHILMLLSQEKLTNTDLAKRLNI
Sbjct 4
              Query 63
Sbjct 64
Query 123
              NKFTDEEOEVISKFLSALTEEFO 145
              F+ EEQ ++ KF++A +EE +
AHFSAEEQVIVEKFITAFSEELE 146
Sbjct 124
>ref|NP_606407.1| d putative repressor protein [Streptococcus pyogenes MGAS8232]
 ref|NP_663873.1|  putative repressor protein [Streptococcus pyogenes MGAS315]
 ref|NP_801332.1|  putative repressor protein [Streptococcus pyogenes SSI-1]
 20 more sequence titles
ref|YP 059443.1|  MarR family transcriptional regulator [Streptococcus pyogenes MGAS10394]
 ref|YP 279544.1| G MarR family transcriptional regulator [Streptococcus pyogenes
MGAS6180]
 ref|YP 595809.1| 🖸 MarR family transcriptional regulator [Streptococcus pyogenes
MGAS9429]
ref|YP 599676.1| Mark family transcriptional regulator (Streptococcus pyogenes
MGAS2096)
 ref|YP 601578.1| G Transcriptional regulator, MarR family [Streptococcus pyogenes
MGAS10750]
 ref|YP 001127671.1| MarR family regulatory protein [Streptococcus pyogenes str. Manfredo]
 ref|YP 002285129.1|  Putative repressor protein [Streptococcus pyogenes NZ131] sp|Q5XEA3.1|ADCR STRP6 RecName: Full=Transcriptional repressor adcR
 gb|AAL96906.1|  putative repressor protein [Streptococcus pyogenes MGAS8232] gb|AAM78676.1|  putative repressor protein [Streptococcus pyogenes MGAS315]
 dbj|BAC63165.1|  putative repressor protein [Streptococcus pyogenes SSI-1]
 gb|AAT86260.1| G Transcriptional regulator, MarR family [Streptococcus pyogenes
MGAS10394]
gb|AAX71189.1|  transcriptional regulator, MarR family [Streptococcus pyogenes MGAS6180]
gb|ABF31265.1|  transcriptional regulator, MarR family (Streptococcus pyogenes MGAS9429)
gb|ABF33144.1| G Transcriptional regulator, MarR family [Streptococcus pyogenes MGAS10270]
gb|ABF37034.1| 🖸 Transcriptional regulator, MarR family [Streptococcus pyogenes
MGAS10750]
 emb|CAM29417.1|  MarR-family regulatory protein [Streptococcus pyogenes str. Manfredo]
 gb[ACI60434.1] G Putative repressor protein [Streptococcus pyogenes NZ131]
Length=147
GENE ID: 994164 adcR | putative repressor protein [Streptococcus pyogenes MGAS8232] (10 or fewer PubMed links)
Score = 141 bits (356), Expect = 1e-32, Method: Compositional matrix adjust. Identities = 69/144 (47%), Positives = 101/144 (70%), Gaps = 0/144 (0%)
              SLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLK 61
+L ++D + TI+ AEN+HE+L G C+SDVKLT+TQEHILMLL++Q TN +A+ L
TLEKKLDNLVNTILLKAENQHELLFGACQSDVKLTNTQEHILMLLSQQRLTNTDLAKALN 62
Ouerv 2
Sbjct 3
               ISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQEL\\ IS AAVTKA+K L +Q+++ ++ T D RV + LTE A P+A EH HH++TL+ Y L\\ ISQAAVTKAIKSLVKQDMLAGTKDTVDARVTYFELTELAKPIASEHTHHHDETLNVYNRL\\ 
Query
        62
                                                                                             121
Sbjct 63
              GNKFTDEEQEVISKFLSALTEEFQ
Query 122
              KF+ +E E++ KF++ EE +
LQKFSAKELEIVDKFVTVFAEELE
Sbjct 123
```

```
subsp. infantarius ATCC BAA-102]
gb|EDT48012.1| hypothetical protein STRINF_00865 [Streptococcus infantarius
subsp. infantarius ATCC BAA-102]
Length=148
  Score = 141 bits (355), Expect = 2e-32, Method: Compositional matrix adjust. Identities = 70/145 (48%), Positives = 103/145 (71%), Gaps = 0/145 (0%)
Query 1
                 MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL
                 M.L. QID + I+ AEN+HE+L G C+S V+LT+TQEHILMLL+++ TN+ +A++L MQLEKQIDCLVNEILLKAENQHELLFGACQSGVELTNTQEHILMLLSQERLTNSALAKRL
Sbjct 1
                 KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
Query 61
                 IS AAVTKA+K L ++ ++ + + +D RV + LTE A PVA EH HH TLS Y++
NISQAAVTKAIKCLVKEGMLAPVKNKDDARVTYFELTEFAKPVADEHNHHHATLSVYKK 120
Sbict 61
Query 121
                 LGNKFTDEEQEVISKFLSALTEEFQ 145
                + + F+DEEQ +IS+FL+A ++E +
MIDDFSDEEQSIISRFLTAFSDELE
Sbict 121
>ref|NP_268489.1| G putative repressor protein [Streptococcus pyogenes M1 GAS]
  ref|YP 281441.1| G MarR family transcriptional regulator [Streptococcus pyogenes
MGAS50051
 gb|AAK33210.1| g putative repressor protein [Streptococcus pyogenes M1 GAS]
gb|AAZ50696.1|  transcriptional regulator, MarR family [Streptococcus pyogenes MGAS5005]
Length=147
GENE ID: 900432 adcR | putative repressor protein [Streptococcus pyogenes M1 GAS] (10 or fewer PubMed links)
 Score = 141 bits (355), Expect = 2e-32, Method: Compositional matrix adjust. Identities = 69/143 (48%), Positives = 100/143 (69%), Gaps = 0/143 (0%)
                 Ouerv 3
                 LEKKLDNLVNTILLKAENQHELLFGACQSDVKLTNTQEHILMLLSQQRLTNTDLAKALNI
Sbjct 4
                 Query 63
Sbict 64
Query 123
                NKFTDEEQEVISKFLSALTEEFQ 145
                KF+ +E E++ KF++ EE +
QKFSAKELEIVDKFVTVFAEELE 146
Sbict 124
>ref|YP 002561480.1| 🖸 MarR-family regulatory protein [Streptococcus uberis 0140J]
 emb|CAR40515.1|  MarR-family regulatory protein [Streptococcus uberis 0140J]
 GENE ID: 7392663 SUB0110 | Mark-family regulatory protein
[Streptococcus uberis 0140J]
 Score = 138 bits (348), Expect = 1e-31, Method: Compositional matrix adjust. Identities = 69/143 (48%), Positives = 102/143 (71%), Gaps = 0/143 (0%)
Query 3
                 LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKI
                                                                                                          62
                L ++IDQ + I+ AEN+HE+L G C+S VKLT+TQEHILMLL+++ TN +A+KL I
LESKIDQLVNQILLKAENQHELLFGACQSHVKLTNTQEHILMLLSQEQLTNTDLAKKLNI
Sbict 4
Query 63
                 SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG
                S AAVTKA+K L + E++ + + T D RV + LT A P+A+EH HH++TL+ Y +L SQAAVTKAIKSLMKHEMLSAIKDTVDARVTYFELTPAAKPIAEEHTQHHDETLNVYTKLL
Sbict 64
Query 123
                NKFTDEEQEVISKFLSALTEEFQ
               + F+ EE+ VI KFL+ ++E +
SSFSSEEKAVIDKFLTVFSDELE 146
Sbict 124
>ref|NP 687190.1| G adc operon repressor AdcR [Streptococcus agalactiae 2603V/R]
 ref[NP_734620.1]  adc operon repressor AdcR [Streptococcus agalactiae NEM316]
 ref[YP 32885.1]  adc operon repressor AdcR [Streptococcus agalactiae A909]
 13 more sequence titles
ref|ZP 00781732.1| repressor protein adcR [Streptococcus agalactiae 18RS21]
ref|ZP 00784062.1| adc operon repressor AdcR [Streptococcus agalactiae H36B]
ref|ZP 00786322.1| adc operon repressor AdcR [Streptococcus agalactiae COH1]
ref|ZP 00790981.1| adc operon repressor AdcR [Streptococcus agalactiae CJB111]
ref|ZP 00790981.1| adc operon repressor AdcR [Streptococcus agalactiae 515]
 gb|AAM99062.1|AE014198 10 🖸 adc operon repressor AdcR [Streptococcus agalactiae 2603V/R]
 emb|CAD45795.1| G unknown [Streptococcus agalactiae NEM316]
 gb|ABA46120.1| G adc operon repressor AdcR [Streptococcus agalactiae A909] repressor protein adcR [Streptococcus agalactiae 18RS21] adc operon repressor AdcR [Streptococcus agalactiae 515] gb|EAO73974.1| adc operon repressor AdcR [Streptococcus agalactiae 515] gb|EAO74935.1| adc operon repressor AdcR [Streptococcus agalactiae CJB111] adc operon repressor AdcR [Streptococcus agalactiae H36B]
 gb | EAO77202.1|
                       adc operon repressor AdcR [Streptococcus agalactiae H36B]
Length=147
```

```
GENE ID: 1012928 adcR | adc operon repressor AdcR [Streptococcus agalactiae 2603V/R] (10 or fewer PubMed links)
 Score = 136 bits (343), Expect = 5e-31, Method: Compositional matrix adjust. Identities = 65/143 (45%), Positives = 99/143 (69%), Gaps = 0/143 (0%)
              Ouerv 3
                                                                                          62
Sbjct 4
              SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG
S AAVTKA+K L Q+++K+++ + D R+ + L+E A P+A EH HH+ TL Y L
Query 63
                                                                                          122
Sbjct 64
              SQAAVTKAVKSLISQDMLKANKDSKDARITYFELSELAKPIADEHTHHHDNTLGVYGRLV
Query 123
              NKFTDEEQEVISKFLSALTEEFQ
                                            145
              N F+ +E+ V+ +FL + E +
NHFSKDEKVVLERFLDLFSRELE
Sbict 124
>ref|YP_001451205.1| G repressor protein adcR [Streptococcus gordonii str. Challis substr.
 qb|AA043167.1|
                    putative transcriptional repressor; AdcR [Streptococcus gordonii
subsp. challis]
 gb|ABV10972.1| 🖸 repressor protein adcR [Streptococcus gordonii str. Challis substr.
CHI)
Length=147
GENE ID: 5599115 adcR | repressor protein adcR [Streptococcus gordonii str. Challis substr. CH1] (10 or fewer PubMed links)
 Score = 133 bits (335), Expect = 4e-30, Method: Compositional matrix adjust. Identities = 75/142 (52%), Positives = 101/142 (71%), Gaps = 0/142 (0%)
Query 3
              LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKI
              LA++ID FL I+ AEN+HEIL+G C S+V LT+TOEHILMLLAE++ TN+ +A+KL + LAHKIDSFLNEIILKAENOHEILVGSCTSNVALTNTOEHILMLLAEEMLTNSDLAKKLNV
Sbjct 4
Query
        63
              SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG
              S AAVTKA+K L \, Q ++++ + D RV + LTE A P+A EH HH TL TYQ L SQAAVTKAVKSLINQGMLETFKDKKDARVTFYRLTELAQPIADEHEHHHAHTLETYQSLL
        64
Sbict
Query 123
              NKFTDEEQEVISKFLSALTEEF
              ++F+ +EQ+ I KFL AL E
DRFSQDEQQAIEKFLEALVGEI
Sbjct 124
>ref[ZP_01817760.1| adc operon repressor AdcR [Streptococcus pneumoniae SP3-BS71]
 700669]
 gb|EDK74363.1|
                    adc operon repressor AdcR [Streptococcus pneumoniae SP3-BS71]
 emb|CAR69937.1|  MarR-family regulatory protein [Streptococcus pneumoniae ATCC
700669]
Length=146
 Score = 132 bits (333), Expect = 6e-30, Method: Compositional matrix adjust. Identities = 71/143 (49%), Positives = 106/143 (74%), Gaps = 0/143 (0%)
               LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKILA ID FL ++ AEN+HEIL+G C S+V LT+TQEHILMLL+E+ TN+++A +L +LAKDIDAFLNEVILQAENQHEILIGHCTSEVALTNTQEHILMLLSEESLTNSELARRLNV \\
Query 3
Sbict 4
              SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG
Query 63
                                                                                          122
                AAVTKA+K L ++ ++++S+ + D RV+ + LT+ A P+A+EH
                                                                       HHE TL TY++
              SQAAVTKAIKSLVKEGMLETSKDSKDARVIFYQLTDLARPIAEEHHHHHEHTLLTYEQVA
Sbjct 64
              NKFTDEEQEVISKFLSALTEEFQ 145
+FT EQ+VI +FL+AL E +
TQFTPNEQKVIQRFLTALVGEIK 146
Ouerv 123
Sbjct 124
pneumonīae SP11-BS70]
 ref|YP 001836857.1| G adc operon repressor AdcR [Streptococcus pneumoniae CGSP14] ref|ZP 02964427.1| putative transcriptional repressor [Streptococcus pneumoniae
CDC0288-04]
 gb|AAL00780.1| G Transcriptional repressor for Zn(2+)-responsive expression [Streptococcus
pneumoniae R6]
gb|EDK63077.1| Transcriptional repressor for Zn(2+)-responsive expression [Streptococcus
pneumoniae SP11-BS70]
 gb|ACB91392.1| G adc operon repressor AdcR [Streptococcus pneumoniae CGSP14]
 gb|EDT94590.1| putative transcriptional repressor [Streptococcus pneumoniae
CDC0288-04]
Length=166
GENE ID: 933868 adcR | adc operon repressor AdcR [Streptococcus pneumoniae R6]
(10 or fewer PubMed links)
Score = 131 bits (329), Expect = 2e-29, Method: Compositional matrix adjust. Identities = 70/143 (48%), Positives = 106/143 (74%), Gaps = 0/143 (0%)
Query 3
              LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKI 62
```

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LA I+ FL ++ AEN+HEIL+G C S+V LT+TQEHILMLL+E+ TN+++A +L + LAKDINAFLNEVILQAENQHEILIGHCTSEVALTNTQEHILMLLSEESLTNSELARRLNV
Sbict 24
               SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG
Query 63
               S AAVTKA+K L ++ ++++S+ + D RV+ + LT+ A P+A+EH HHE TL TY+++
SOAAVTKAIKSLVKEGMLETSKDSKDARVIFYOLTDLARPIAEEHHHHHEHTLLTYEOVA
Sbict 84
Query 123
               NKFTDEEQEVISKFLSALTEEFQ 145
               +FT EQ+VI +FL+AL E +
TQFTPNEQKVIQRFLTALVGEIK 166
Sbict 144
>ref[NP 346586.1] G adc operon repressor AdcR [Streptococcus pneumoniae TIGR4]
 ref[ZP 01408975.1]  hypothetical protein SpneT 02000553 [Streptococcus pneumoniae
TIGR4]
 ref[YP 817386.1|  adc operon repressor AdcR [Streptococcus pneumoniae D39]
 29 more sequence titles
ref|ZP 01820229.1|
ref|ZP 01821749.1|
ref|ZP 01827937.1|
ref|ZP 01830531.1|
ref|ZP 01830186.1|
ref|ZP 0183060.1|
                         adc operon repressor AdcR [Streptococcus pneumoniae SP6-BS73] adc operon repressor AdcR [Streptococcus pneumoniae SP9-BS68]
                          adc operon repressor AdcR [Streptococcus pneumoniae SP14-BS69] adc operon repressor AdcR [Streptococcus pneumoniae SP18-BS74] adc operon repressor AdcR [Streptococcus pneumoniae SP18-BS74] adc operon repressor AdcR [Streptococcus pneumoniae SP19-BS75] adc operon repressor AdcR [Streptococcus pneumoniae SP23-BS72]
ref[ZP_02708953.1|
CDC1873=00]
                          putative transcriptional repressor [Streptococcus pneumoniae
ref[ZP 02710784.1|
CDC1087-00]
                          putative transcriptional repressor [Streptococcus pneumoniae
 ref|ZP_02713441.1|
                           putative transcriptional repressor [Streptococcus pneumoniae
SP1951
ref|ZP 02718213.1|
CDC3059=06]
                          putative transcriptional repressor [Streptococcus pneumoniae
 ref|ZP 02721950.1|
                          putative transcriptional repressor [Streptococcus pneumoniae
MLV-016T
 ref|YP 001695528.1| 💆 putative transcriptional repressor [Streptococcus pneumoniae
HungaryT9A-6]
 ref|YP 002038763.1|  transcriptional regulator, MarR family [Streptococcus pneumoniae
G541
 emb|CAA96184.1|
                      _AdcR protein [Streptococcus pneumoniae]
 gb|AAK76226.1| G adc operon repressor AdcR [Streptococcus pneumoniae TIGR4]
 gb | EDK81745.1 |
                     adc operon repressor AdcR [Streptococcus pneumoniae SP23-BS72]
 gb|ACA36761.1| G putative transcriptional repressor [Streptococcus pneumoniae
Hungary19A-6]
gb[EDT50754.1|
CDC1873-00]
                     putative transcriptional repressor [Streptococcus pneumoniae
gb|EDT91227.1|
CDC1087-00]
                     putative transcriptional repressor [Streptococcus pneumoniae
gb[EDT92758.1|
SP195]
                     putative transcriptional repressor [Streptococcus pneumoniae
 gb|EDT96372.1|
                     putative transcriptional repressor [Streptococcus pneumoniae
CDC3059-06]
 gb|EDT98576.1|
                     putative transcriptional repressor [Streptococcus pneumoniae
MLV-0161
gb|ACF56415.1|  transcriptional regulator, MarR family (Streptococcus pneumoniae G54)
Length=146
GENE ID: 931929 SP 2172 | adc operon repressor AdcR [Streptococcus pneumoniae TIGR4] (10 or fewer PubMed links)
 Score = 130 bits (328), Expect = 2e-29, Method: Compositional matrix adjust. Identities = 70/143 (48%), Positives = 106/143 (74%), Gaps = 0/143 (0%)
               LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKI
LA I+ FL ++ AEN+HEIL+G C S+V LT+TQEHILMLL+E+ TN+++A +L +
LAKDINAFLNEVILQAENQHEILIGHCTSEVALTNTQEHŢLMLLSEESLTNSELARRLNV
Query 3
Sbjct 4
Query 63
               SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG
               S AAVTKA+K L ++ ++++S+ + D RV+ + LT+ A P+A+EH HHE TL TY+++
SQAAVTKAIKSLVKEGMLETSKDSKDARVIFYQLTDLARPIAEEHHHHHEHTLLTYEQVA
Sbict 64
Query 123
               NKFTDEEQEVISKFLSALTEEFQ
               +FT EQ+VI +FL+AL E +
TQFTPNEQKVIQRFLTALVGEIK 146
Sbict 124
>ref|YP 138725.1| 🖸 zinc transport transcriptional repressor [Streptococcus thermophilus
LMG 183T1]
 ref|YP 140614.1| 🖸 zinc transport transcriptional repressor [Streptococcus thermophilus
CNRZ1066)
 qb|AAV59910.1| G zinc transport transcriptional repressor [Streptococcus thermophilus
LMG 183111
 gb|AAV61799.1| C zinc transport transcriptional repressor [Streptococcus thermophilus
CNRZ10661
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GENE ID: 3164452 adcR | zinc transport transcriptional repressor [Streptococcus thermophilus LMG 18311] (10 or fewer PubMed links)
 Score = 129 bits (325), Expect = 5e-29, Method: Compositional matrix adjust. Identities = 62/145 (42%), Positives = 104/145 (71%), Gaps = 0/145 (0%)
Query 1
                  MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL
                  + L Q++Q + I+ AEN++E+L+G+C S VKLT+TQEHILMLL+E TN+++A+ L IELEEQVNQLINQILLKAENQYELLIGQCRSKVKLTNTQEHILMLLSEGOKTNSELAKAL
Sbjct 6
Query 61
                  KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
                  +S AAVTKA+K L ++ +++ + + D RV + LT++A P+A+EH HH++TL Y+
NVSQAAVTKAVKTLVKEGMLEGKKDKDDGRVTYFVLTQEAQPIAQEHKEHHQETLGVYRS
Sbjct
          66
Query 121
                  LGNKFTDEEQEVISKFLSALTEEFQ 145
                  + ++F +E++VI +FL L E+ +
VLDQFDHQERQVIGRFLIKLAEKIE
Sbjct 126
>ref|YP 819746.1| 🗲 zinc transport transcriptional repressor [Streptococcus thermophilus
  gb|ABJ65550.1| G transcriptional regulator, MarR family [Streptococcus thermophilus
LMD-91
Length=147
GENE ID: 4438531 STER 0233 | zinc transport transcriptional repressor [Streptococcus thermophilus LMD-9] (10 or fewer PubMed links)
 Score = 128 bits (322), Expect = 1e-28, Method: Compositional matrix adjust. Identities = 61/145 (42%), Positives = 104/145 (71%), Gaps = 0/145 (0%)
Query 1
                  {\tt MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL}
                  + L +++Q + I+ AEN++E+L+G+C S VKLT+TQEHILMLHE TN+++A+ L IELEERVNQLINQILLKAENQYELLIGQCRSKVKLTNTQEHILMLLSEGQKTNSELAKAL
Sbjct 2
Query 61
                  KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
                  +S AAVTKA+K L ++ +++ + +D RV + LT++A P+A+EH HH++TL Y+
NVSQAAVTKAVKTLVKEGMLEGKKDKDDGRVTYFVLTQEAQPIAQEHKEHHQETLGVYRS
Sbjct 62
                  LGNKFTDEEQEVISKFLSALTEEFQ 145
Query 121
                  + ++F +E++VI +FL L E+ +
VLDQFDHQERQVIGRFLIKLAEKIE
Sbjct 122
>ref|YP_001197479_1| G transcriptional regulator [Streptococcus suis 05ZYH33]
 gb|ABP89079.1| G Transcriptional regulator [Streptococcus suis 05ZYH33]
Length=149
GENE ID: 5099418 SSU05 0109 | transcriptional regulator [Streptococcus suis 052\text{YH33}] (10 or fewer PubMed links)
 Score = 116 bits (290), Expect = 7e-25, Method: Compositional matrix adjust. Identities = 70/143 (48%), Positives = 100/143 (69%), Gaps = 0/143 (0%)
                   \begin{array}{llll} LANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKI\\ +A&+I+++L&I+&+EN+&EIL+G&C+S&VKLT+TQEHILML+&+&TN&+IA++L&+\\ IALEIEKYLHEIVLSSENQLEILVGSCQSTVKLTNTQEHILMLIEKAAYTNTEIAKELNV \end{array}
Ouerv 3
Sbjct 6
                  SPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELG S AA+TKA K L Q L+ + R D R+V +SLTE A P+A EHA HH TL Y+EL SQAAITKATKSLVAQGLLVAVRDDKDARIVRFSLTEAAKPIAAEHAHHHAHTLEAYEELL
Query 63
Sbjct 66
                  NKFTDEEQEVISKFLSALTEEFQ
Ouerv 123
                 ++ EEQE I++FLS L E+ +
ENYSLEEQESIARFLSELVEKIR 148
Sbjct 126
>ref|ZP 03625066.1| transcriptional regulator, MarR family [Streptococcus suis 89/1591]
 gb|EEF64628.1| transcriptional regulator, Mark family [Streptococcus suis 89/1591]
Length=149
 Score = 110 bits (275), Expect = 4e-23, Method: Compositional matrix adjust. Identities = 69/140 (49%), Positives = 98/140 (70%), Gaps = 0/140 (0%)
Ouerv 6
                 QIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKISPA +I+++L I+ +EN+ EIL+G C+S VKLT+TQEHILML+ + TN +IA++L +S A
Sbjct 9
                  EIEKYLHEIVLSSENQLEILVGSCQSTVKLTNTQEHILMLIEKAAYTNTEIAKELNVSQA
                 AVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKF A+TKA K L Q L+ + R D R+V +SLTE A P+A EHA HH TL Y+EL + AITKATKSLVAQGLLVAVRDDKDARIVRFSLTEAAKPIATEHAHHHAHTLEAYEELLEHY
Ouerv 66
                                                                                                                 125
Sbjct 69
Ouerv 126
                 TDEEOEVISKFLSALTEEFO 145
Sbjct 129
                 SLEEQESIARFLSELVEKIR 148
>emb|CAA75313.1| hypothetical protein [Lactococcus lactis subsp. cremoris]
Length=48
 Score = 92.4 bits (228), Expect = 1e-17, Method: Compositional matrix adjust. Identities = 43/47 (91%), Positives = 46/47 (97%), Gaps = 0/47 (0%)
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MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLA 47
MSLANQIDQFLG IMQFAENKHEILLG+CES+VKLTSTQEHILM+LA MSLANQIDQFLGAIMQFAENKHEILLGECESNVKLTSTQEHILMILA 47
Query 1
 Sbjct 1
>ref[ZP_00365496.1| COG1846: Transcriptional regulators [Streptococcus pyogenes M49
 5911
Length=103
  Score = 90.9 bits (224), Expect = 3e-17, Method: Compositional matrix adjust. Identities = 45/102 (44%), Positives = 68/102 (66%), Gaps = 0/102 (0%)
                  MLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPV
                 MLL++Q TN +A+ L IS AAVTKA+K L +Q+++ ++ T D RV + LTE A P+
MLLSQQRLTNTDLAKALNISQAAVTKAIKSLVKQDMLAGTKDTVDARVTYFELTELÄKPI
Sbict 1
                 AKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ 145
A EH HH++TL+ Y L KF+ +E E++ KF++ EE +
ASEHTHHHDETLNVYNRLLQKFSAKELEIVDKFVTVFAEELE 102
Query 104
Sbjct 61
gb|ABV60899.1| G possible MarR family transcriptional regulator [Bacillus pumilus
SAFR-032]
gb|EDW20313.1| YvnA [Bacillus pumilus ATCC 7061]
Length=152
GENE ID: 5619427 BPUM 0200 | MarR family transcriptional regulator [Bacillus pumilus SAFR-032] (10 or fewer PubMed links)
 Score = 67.0 bits (162), Expect = 5e-10, Method: Compositional matrix adjust. Identities = 40/111 (36%), Positives = 66/111 (59%), Gaps = 2/111 (1%)
                 Query 36
Sbjct 42
                 WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF
+SLTEK +A H HEK +++Y E +F ++E +VI +FL A E+
YSLTEKGTQLAALHDELHEKAVASYLEFLQQFHEDELQVIERFLKAWKEKI
Ouerv 94
Sbict 102
>ref|YP_001199672.1|  transcriptional regulator [Streptococcus suis 98HAH33] gb|ABP91272.1|  Transcriptional regulator [Streptococcus suis 98HAH33] Length=87
GENE ID: 5101832 SSU98 0112 | transcriptional regulator [Streptococcus suis 98HAH33] (10 or fewer PubMed links)
 Score = 63.2 bits (152), Expect = 6e-09, Method: Compositional matrix adjust. Identities = 41/86 (47%), Positives = 56/86 (65%), Gaps = 0/86 (0%)
                 LKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQ 119 + +S AA+TKA K L Q L+ + R D R+V +SLTE A P+A EHA HH TL Y+ MNVSQAAITKATKSLVAQGLLVAVRDDKDARIVRFSLTEAAKPIAAEHAHHHAHTLEAYE 60
Ouerv 60
Sbjct 1
                 ELGNKFTDEEQEVISKFLSALTEEFQ 145
EL ++ EEQE I++FLS L E+ +
ELLENYSLEEQESIARFLSELVEKIR 86
Query 120
Sbjct 61
>ref|YP 187318.1|  MarR family transcriptional regulator [Staphylococcus aureus
subsp. aureus COL]
 ref|YP_495086.1|  MarR family transcriptional regulator [Staphylococcus aureus
subsp. aureus USA300]
ref|YP_001576367.1| G MarR family transcriptional regulator [Staphylococcus aureus subsp. aureus USA300_TCH1516]
 ref|ZP 02761767.1| G MarR family transcriptional regulator [Staphylococcus aureus
subsp. aureus USA300_TCH1516]
 gb|AAW37302.1| C transcriptional regulator, MarR family [Staphylococcus aureus
subsp. aureus COL
gb|ABD21126.1| G transcriptional regulator, MarR family [Staphylococcus aureus subsp. aureus USA300_FPR3757]
 gb|ABX30488.1| G MarR family transcriptional regulator [Staphylococcus aureus
subsp. aureus USA300 TCH1516]
Length=154
GENE ID: 3238188 SACOL2524 | Mark family transcriptional regulator [Staphylococcus aureus subsp. aureus COL] (10 or fewer PubMed links)
 Score = 63.2 bits (152), Expect = 7e-09, Method: Compositional matrix adjust. Identities = 40/112 (35%), Positives = 66/112 (58%), Gaps = 1/112 (0%)
                 LTSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93
L+ TQ HI+ L+ N K ++E L +S A+TK++KKL ++L+ S ++R V
LSLTQFHIIELIDNNDKVNNKFLSEMLNVSKPAITKSIKKLLAKDLVVESHNEFNKREVN 95
Ouerv 35
Sbjct 36
                 WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ
Ouerv 94
                 +SLT+K ++ H HEK++ Y+E+ F D+E VI +FL+ EE +
YSLTQKGKKLSYIHDELHEKSVKKYEEVLKVFDDDEMAVIIEFLNRSIEELK
Sbict 96
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>ref|YP 001727325.1| G transcriptional repressor for Zn(2+)-responsive expression [Leuconostoc
citreum KM201
 gb|ACA81881.1|  Transcriptional repressor for Zn(2+)-responsive expression [Leuconostoc
citreum KM20]
Length=146
GENE ID: 6063203 adcR | transcriptional repressor for Zn(2+)-responsive expression [Leuconostoc citreum KM20] (10 or fewer PubMed links)
 Score = 62.8 bits (151), Expect = 8e-09, Method: Compositional matrix adjust. Identities = 41/138 (29%), Positives = 72/138 (52%), Gaps = 2/138 (1%)
Ouerv 1
               {\tt MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL}
               MS ++ I Q L T +Q E + + + K+ +TQ H+LMLL Q +TN+ +AE + MSQSDHIIQELNTFVQTYAASSEFI--QTTAAQKINATQAHLLMLLKTQHATNSSLAESM
Sbjct 1
Query 61
               KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE 120 ++ A+TKA+K L + +++ ND+R V + L+ + +A +H H
               HLTKPAITKAIKNLIAHGYVVATKDVNDKRSVNYQLSTEGMQLAAQHEASHRNLHHRIDH 118
Sbjct 59
Query 121
               LGNKFTDEEQEVISKFLS 138
               FT ++E I FL+
TIATFTPAQRETIVAFLA
Sbict 119
ref|YP_001333445.1| 🖸 transcriptional regulator MarR family protein (Staphylococcus
aureus NCTC 8325]
 dbj|BAF68683.1|  transcriptional regulator MarR family protein [Staphylococcus
aureus subsp. aureus str. Newman]
Length=152
 GENE ID: 3921258 SAOUHSC_02819 | hypothetical protein
[Staphylococcus aureus subsp. aureus NCTC 8325]
 Score = 62.8 bits (151), Expect = 8e-09, Method: Compositional matrix adjust. Identities = 40/112 (35%), Positives = 66/112 (58%), Gaps = 1/112 (0%)
               LTSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93
L+ TQ HI+ L+ N K ++E L +S A+TK++KKL ++L+ S ++R V
LSLTQFHIIELIDNNDKVNNKFLSEMLNVSKPAITKSIKKLLAKDLVVESHNEFNKREVN 93
Query 35
Sbjct 34
               WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ 145
+SLT+K ++ H HEK++ Y+E+ F D+E VI +FL+ EE +
YSLTQKGKKLSYIHDELHEKSVKKYEEVLKVFDDDEMAVIIEFLNRSIEELK 145
Query 94
Sbict 94
>ref|ZP_03563431.1| MarR family transcriptional regulator [Staphylococcus aureus
subsp. aureus str. JKD6008]
ref|ZP_03566444.1| MarR family transcriptional regulator [Staphylococcus aureus
subsp. aureus str. JKD6009]
Length=154
 Score = 61.2 bits (147), Expect = 3e-08, Method: Compositional matrix adjust. Identities = 40/112 (35%), Positives = 65/112 (58%), Gaps = 1/112 (0%)
Query 35
               LTSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93
L+ TQ HI+ L+ N K ++E L +S A+TK++KKL ++L+ S ++R V
LSLTQFHIIELIDNNDKVNNKFLSEMLNVSKPAITKSIKKLLAKDLVVESHNEFNKREVN 95
Sbjct 36
               Ouerv 94
Sbjct 96
>ref[YP_174450.1]  MarR family transcriptional regulator [Bacillus clausii KSM-K16]
 dbj|BAD63489.1|  MarR family transcriptional regulator [Bacillus clausii KSM-K16]
Length=153
 GENE ID: 3204362 ABC0950 | MarR family transcriptional regulator
[Bacillus clausii KSM-K16] (10 or fewer PubMed links)
 Score = 60.5 bits (145), Expect = 4e-08, Method: Compositional matrix adjust. Identities = 40/128 (31%), Positives = 73/128 (57%), Gaps = 3/128 (2%)
               AENKHEILLGKCESDV--KLTSTQEHILMLLAEQ-ISTNAKIAEKLKISPAAVTKALKKL 74 A+ + + + G E + T TQ HI+ ++ EQ + N +AE L +S A+TKA+KKL ADRRRKAMKGSQEESIVSDWTLTQLHIVAIVKEQERANNTMLAEHLNVSKPAITKAVKKL 80
Ouerv 18
Sbjct 21
               Ouerv 75
               LDQQILEKTQQADNKKEVYYRLTKSGEMLAFIHSQLHEQARNRYMRIFAEFNSTELETII 140
Sbjct 81
               KFLSALTE
Query 135
Sbict 141
              RFLHALAE 148
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13
                                                                                                                    Annex 1
>ref|YP 080661.1 G transcriptional regulator YvnA [Bacillus licheniformis ATCC 14580]
 gb|AAU25023.1| G probable transcriptional regulator YvnA [Bacillus licheniformis
ATCC 14580]
Length=157
GENE ID: 3028745 yvnA | transcriptional regulator YvnA [Bacillus licheniformis ATCC 14580] (10 or fewer PubMed links)
 Score = 59.3 bits (142), Expect = 1e-07, Method: Compositional matrix adjust. Identities = 37/105 (35%), Positives = 64/105 (60%), Gaps = 2/105 (1%)
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TSTQEHILMLLAEQIS~~TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93 Ouerv 36 T TQ HI+ L++E + NA +A KL+IS AAVTKAY L + +I+S + N+ + +
TLTQLHIISLISESEADVNNAFLAAKLQISKAAVTKAVNVLTKHGMIESHKKPNNNKELY 106 Sbjct 47 WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS 138 ++LT++ +A H HE Y EL ++F++ E + + +FL+ Query 94

Sbict 107 YTLTDEGKKLADIHDRMHEIAKQRYIELFDRFSESELQTVIRFLN

>ref|YP 254603.1| 🖸 hypothetical protein pSHaeC05 [Staphylococcus haemolyticus JCSC1435] dbj|BAE05997.1| G unnamed protein product [Staphylococcus haemolyticus JCSC1435] Length=155

GENE ID: 3431756 pSHaeC05 | hypothetical protein
[Staphylococcus haemolyticus JCSC1435] (10 or fewer PubMed links)

Score = 59.3 bits (142), Expect = 1e-07, Method: Compositional matrix adjust. Identities = 38/119 (31%), Positives = 69/119 (57%), Gaps = 1/119 (0%)

KCESDVKLTSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRAT 86 K D+ L+ TO HI+ ++ + N K +AE+L +S AVTK++KKL +EL+ Query 28 Sbjct 29 KGNEDMDLSLTQFHIIEIIDKHEKVNNKFLAEELNVSKPAVTKSIKKLLSKELVVELNNE

Query 87 NDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ

+++R V ++LT++ ++ H H+K + Y+E+ F ++E E I +FL +E +
SNKREVYYNLTKRGEKLSFIHDDLHKKAVKKYEEVLKVFDEKEMETIIEFLKRSVDELK Sbjct 89

>ref|YP 093087.1| G YvnA [Bacillus licheniformis ATCC 14580] gb|AAU42394.1| G YvnA [Bacillus licheniformis DSM 13] Length=160

GENE ID: 3100261 yvnA | similar to proteins from B. subtilis [Bacillus licheniformis ATCC 14580] (10 or fewer PubMed links)

Score = 59.3 bits (142), Expect = 1e-07, Method: Compositional matrix adjust. Identities = 37/105 (35%), Positives = 64/105 (60%), Gaps = 2/105 (1%)

Query 36 TSTQEHILMLLAEQIS--TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93 T TQ HI+ L++E + NA +A KL+IS AAVTKA+ L + +I+S + N+ + + TLTQLHIISLISESEADVNNAFLAAKLQISKAAVTKAVNVLTKHGMIESHKKPNNNKELY 109 Sbjct 50

WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS 138 ++LT++ +A H HE Y EL ++F++ E + + +FL+ Query 94 YTLTDEGKKLADIHDRMHEIAKQRYIELFDRFSESELQTVIRFLN 154 Sbict 110

>ref|ZP\_00738634.1| Transcriptional regulator, MarR family [Bacillus thuringiensis serovar israelensis ATCC 35646] gb|EAO57135.1| Transcriptional regulator, MarR family [Bacillus thuringiensis serovar israelensis ATCC 35646] Length=156

Score = 55.8 bits (133), Expect = 9e-07, Method: Compositional matrix adjust. Identities = 32/106 (30%), Positives = 64/106 (60%), Gaps = 2/106 (1%)

Query 36 Sbict 46

Query 94 WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSA 139 ++LT+K + +A H H+ Y EL +F D E ++++KFL A
YTLTDKGIQLAIVHKKLHKIAHERYSELFQQFNDSELQIVTKFLEA 151 Sbjct 106

>ref|ZP 02613091.1| transcriptional regulator, MarR family [Clostridium botulinum NCTC 29T6] gb|EDT83321.1| transcriptional regulator, MarR family [Clostridium botulinum Length=174

Score = 54.3 bits (129), Expect = 3e-06, Method: Compositional matrix adjust. Identities = 35/113 (30%), Positives = 62/113 (54%), Gaps = 2/113 (1%)

ESDVKLTSTQE-HILMLLAEQ-ISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATN ++D+K S E H++ + + + S N IA +L ++ ++K KL +++IK+ + N DNDIKGISLSEFHVIECIGKNNMSNNIFIARELNMTKGGISKINSKLLSKDIIKADKIEN Query 30

Sbjct 58

DERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL Query 88 D+R + +SLTEK + + K H HEK ++ + + EE I KFL L
DKREIYYSLTEKGIALFKLHEYLHEKEREKLMKILSNYKLEEITTILKFLEDL Sbjct 118

14 Annex 1 >ref|YP 001253547.1| 🖸 MarR family transcriptional regulator [Clostridium botulinum A str. ATCC 3502] ref|YP 001386937.1| G MarR family transcriptional regulator [Clostridium botulinum A str. Hall] emb|CAL82569.1| MarR-family transcriptional regulator {Clostridium botulinum A str. ATCC 3502] gb|ABS35317.1| G transcriptional regulator, MarR family [Clostridium botulinum A str. ATCC 19397] gb|ABS36227.1| 🖸 transcriptional regulator, MarR family [Clostridium botulinum A str. Hall] Length=174 GENE ID: 5185271 CBO1016 | MarR family transcriptional regulator [Clostridium botulinum A str. ATCC 3502] (10 or fewer PubMed links) Score = 54.3 bits (129), Expect = 3e-06, Method: Compositional matrix adjust. Identities = 35/113 (30%), Positives = 63/113 (55%), Gaps = 2/113 (1%) ESDVKLTSTQE-HILMLLAEQ-ISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATN 87 ++D+K S E H++ + + +S N IA++L ++ ++K KL +++IK+ + N DNDIKGISLSEFHVIECIGKNNMSNNIFIAKELNMTKGGISKINSKLLSKDIIKADKIEN 117 Query 30 Sbjct 58 Query 88 DERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL 140 D+R + +SLTEK + + K H HEK ++ + + EE I KFL L
DKREIYYSLTEKGIALFKLHEHLHEKEREKLMKILSNYKLEEITTILKFLEDL 170 Sbjct 118 >ref|YP\_001780642.1| G MarR family transcriptional regulator [Clostridium botulinum Bl str. Okral gb|ACA44681.1| 🖸 transcriptional regulator, MarR family [Clostridium botulinum Bl str. Okra] Length=174 GENE ID: 6149223 CLD\_3550 | MarR family transcriptional regulator [Clostridium botulinum B1 str. Okra] (10 or fewer PubMed links) Score = 54.3 bits (129), Expect = 3e-06, Method: Compositional matrix adjust. Identities = 35/113 (30%), Positives = 63/113 (55%), Gaps = 2/113 (1%) Query 30 Sbjct 58 Query 88 DERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL 140 D+R + +SLTEK + + K H HEK ++ + + EE I KFL L
DKREIYYSLTEKGIALFKLHEHLHEKEREKLMKILSNYKLEEITTILKFLEDL Sbict 118 cremoris MG1363] emb|CAL98430.1| transcriptional regulator, MarR family [Lactococcus lactis subsp. cremoris MG1363] Length=295 GENE ID: 4797387 rmaB | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links) Score = 53.9 bits (128), Expect = 4e-06, Method: Compositional matrix adjust. Identities = 38/101 (37%), Positives = 54/101 (53%), Gaps = 3/101 (2%) Query 42 ILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAV 101 ++ L E TNA+IAE L I P++VT +K+LEEE++ + ND+RV LTEK
LVELWNEDGLTNAEIAELLDIKPSSVTTQVKQLEEAEMVIRKQDENDKRVNRIFLTEKGR 106 Sbjct 47 PVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTE 142 Ouerv 102 + T H T GN TDEEQE ++ + L E EAQETROTMHNDISETI--FGN-LTDEEQEQLANLMEKLVE 144 >ref|ZP\_02617530.1| transcriptional regulator, MarR family [Clostridium botulinum gb|EDT85893.1| transcriptional regulator, MarR family [Clostridium botulinum Bf] Length=174 Score = 53.9 bits (128), Expect = 4e-06, Method: Compositional matrix adjust. Identities = 35/113 (30%), Positives = 62/113 (54%), Gaps = 2/113 (1%) ESDVKLTSTQE-HILMLLAEQ-ISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATN 87 ++D+K S E H++ + + +S N IA+ L ++ ++K KL +++IK+ + N DNDIKGISLSEFHVIECIGKNNMSNNIFIAKDLNMTKGGISKINSKLLSKDIIKADKIEN 117 Query 30

Query 88 DERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL 140 D+R + + SLTEK + + K H HEK ++ + + EE I KFL L
DKREIYYSLTEKGIALFKLHEHIHEKEREKLMKILSNYKLEEITTILKFLEDL 170 Sbjct 118

Sbjct 58

>ref|YP 861960.1| G MarR family transcriptional regulator [Gramella forsetii KT0803] emb|CAL66893.1| MarR family transcriptional regulator protein [Gramella forsetii

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Annex 1
KT0803]
Length=158
GENE ID: 4650237 GFO 1928 | MarR family transcriptional regulator [Gramella forsetii KT\overline{0}803] (10 or fewer PubMed links)
 Score = 53.9 bits (128), Expect = 4e-06, Method: Compositional matrix adjust. Identities = 39/112 (34%), Positives = 65/112 (58%), Gaps = 8/112 (7%)
                MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQ--ISTNAKIAE
+ L NQI + ++ + ++ L K E LT Q +L++L E+ +S N KI E
LKLENQICFPIYSVSRLITKAYKPYLDKLE----LTYPQYLVLLVLWEEHKLSVN-KIGE
Ouerv 1
Sbjct 7
                KLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATH 110 KL ++ ++ LK++++ EL+K +R++NDER VL LT+K + KE A H KLMLNTNTLSPLLKRMEKNELLKRNRSSNDERTVLVGLTDKGLSY-KEKAAH 112
Ouerv 59
Sbjct 62
>ref[NP 391388.1] G hypothetical protein BSU35080 (Bacillus subtilis subsp. subtilis
str. 168]
ref[ZP_03593305.1] hypothetical protein Bsubsl_18986 [Bacillus subtilis subsp. subtilis
str. 168]
ref|ZP_03597590.1| hypothetical protein BsubsN3_18902 [Bacillus subtilis subsp. subtilis str. NCIB 3610]
 6 more sequence titles
ref[ZP_03601994.1] hypothetical protein BsubsJ_18865 [Bacillus subtilis subsp. subtilis
str. JH642)
ref[ZP 03606279.1] hypothetical protein BsubsS_19021 [Bacillus subtilis subsp. subtilis
 sp|P40762.1|YVMB BACSU RecName: Full=Uncharacterized HTH-type transcriptional regulator
 emb|CAA85355.1| unnamed protein product [Bacillus subtilis] gb|AAC67278.1| YzhA [Bacillus subtilis]
 Length=169
GENE ID: 936625 vvmB | vvmB [Bacillus subtilis subsp. subtilis str. 168]
(10 or fewer PubMed links)
 Score = 53.5 bits (127), Expect = 5e-06, Method: Compositional matrix adjust. Identities = 36/91 (39%), Positives = 53/91 (58%), Gaps = 2/91 (2%)
                TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEH-ATH 110 NA IA K+ +S A VTK KL ++E I S + T++++ V + LT K + H H NNAGIARKMNLSKANVTKISTKLIKEEFINSYQLTDNKKEVYFKLTRKGRRIFDLHEKLH 122
Query 52
Sbjct 63
Query 111
                HEKTLSTYQELGNKFTDEEQEVISKFLSALT
+K L+ YQ L + F+ EEQ+ + KFL LT
Sbjct 123 KKKELAFYQFL-DSFSQEEQKAVLKFLEQLT
>ref[ZP_02994615.1| hypothetical protein CLOSPO_01734 [Clostridium sporogenes ATCC
gb(EDU38872.1| hypothetical protein CLOSPO_01734 (Clostridium sporogenes ATCC
15579)
Length=174
 Score = 53.5 bits (127), Expect = 6e-06, Method: Compositional matrix adjust. Identities = 34/113 (30%), Positives = 63/113 (55%), Gaps = 2/113 (1%)
                 Query 30
Sbjct 58
                 DERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL 140 D+R + +SLTEK + + K H H+K ++ + EE I KFL L
Query 88
Sbjct 118
                DKREIYYSLTEKGIVLFKLHEYLHKKEQEKLMKILSNYKQEEITTILKFLDDL 170
>ref|ZP 01994138.1| hypothetical protein DORLON 00120 [Dorea longicatena DSM 13814]
gb|EDM64274.1| hypothetical protein DORLON_001Z0 [Dorea longicatena DSM 13814]
Length=153
 Score = 53.1 bits (126), Expect = 6e-06, Method: Compositional matrix adjust. Identities = 36/111 (3%), Positives = 55/111 (49%), Gaps = 4/111 (3%)
                LTSTQEHILMLLAEQISTN-AKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93
LT+ H++ + N + IA KL I+ ++T A+ L + ++ R+ D RVVL
LTNNDMHVIEAVGLGDGNNMSSIARKLNITVGSLTTAMNSLVNKRYVERHRSEEDRRVVL 96
Query 35
Sbjct 37
                 {\tt WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF}
Query 94
                 LTEK V H +H + Q + +K D E V+ K L AL+E F
VKLTEKGVKAYHHHEDYHRQMT---QAILDKLDDTELPVLVKTLDALSEFF 144
Sbjct 97
>ref|ZP 03294064.1| hypothetical protein CLOHIR_02015 [Clostridium hiranonis DSM
```

13275] gb|EÉA84316.1| hypothetical protein CLOHIR\_02015 [Clostridium hiranonis DSM 13275] Length=166

Score = 53.1 bits (126), Expect = 7e-06, Method: Compositional matrix adjust. Identities = 37/131 (28%), Positives = 68/131 (51%), Gaps = 6/131 (4%)

TIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTN-AKIAEKLKISPAAVTKAL 71

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T +F HE G+ K+ + H++ + E N ++++E+L I+ AV++ L
TYYKFLSTPHEYYPGE----KMHMREVHVITEIGEGGLDNISELSERLNITKGAVSQYL 74
Sbjct 20
                KKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQE 131
Ouerv
         72
                KKL+++ I+ + + D+R LTEK + K H + E+ + N+FT+EE E
KKLEKKGFIERVQESEDKRQYSVRLTEKGKELDKIHTKYDEEQYAKACPFFNEFTEEELE
Sbjct
         75
Query 132
                VISKFLSALTE 142
                LICRFEARFAE 145
Sbjct 135
>ref|ZP 02211326.1| hypothetical protein CLOBAR 00939 [Clostridium bartlettii DSM
 gb|EDQ97187.1| hypothetical protein CLOBAR_00939 [Clostridium bartlettii DSM
Length=150
 Score = 52.4 bits (124), Expect = 1e-05, Method: Compositional matrix adjust. Identities = 33/94 (35%), Positives = 50/94 (53%), Gaps = 3/94 (3%)
                TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHH 111 T +IA L+I+ +T A+ +L ++ + SR D RVVL SLTEK K HA H
Ouerv 52
                T +IA L+I+ +T A+ +L ++ + SR D RVVL SLTEK K HA H
TMGEIAHDLRITVGTLTSAINRLIKKGYAERSRTEEDRRVVLVSLTEKGKHAYKIHADFH 112
Sbjct 53
                EKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ
Query 112
++ + Q N + DEEQEV+ + + F+
Sbjct 113 KEMV---QATLNSYNDEEQEVLCDVIEKINIFFE 143
gb|ABS39442.1|  transcriptional regulator, MarR family [Clostridium botulinum F str. Lange]and]
  str. Langeland]
Length=174
GENE ID: 5405444 CLI 1098 | MarR family transcriptional regulator [Clostridium botulinum F str. Langeland]
 Score = 52.4 bits (124), Expect = 1e-05, Method: Compositional matrix adjust. Identities = 34/113 (30%), Positives = 62/113 (54%), Gaps = 2/113 (1%)
Ouerv 30
               ESDVKLTSTQE-HILMLLAEQ-ISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATN 87 ++D+K S E H++ + + + S N IA++L ++ ++K KL +++IK+ + N
                DNDIKGISLSEFHVIECIGKNNMSNNIFIAKELNMTKGGISKINSKLLSKDIIKADKIEN 117
Sbjct 58
                DERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL
Query 88
               D+R + +SLTEK + + K H HEK ++ + + EE I FL L
DKREIYYSLTEKGIALFKLHEHLHEKEREKLMKILSNYKLEEITTILNFLEDL
gb|ACA55750.1| G transcriptional regulator, MarR family [Clostridium botulinum A3 str. Loch Maree]
Length=174
GENE ID: 6154364 CLK 0459 | MarR family transcriptional regulator [Clostridium botulinu\overline{m} A3 str. Loch Maree] (10 or fewer PubMed links)
 Score = 51.6 bits (122), Expect = 2e-05, Method: Compositional matrix adjust. Identities = 33/113 (29%), Positives = 62/113 (54%), Gaps = 2/113 (1%)
               ESDVKLTSTQE-HILMLLAEQ-ISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATN 87 ++D+K S E H++ + + + + N 'IA++L ++ ++K KL +++I++ + N
Query 30
                ++D+K S E H++ + + + N 'IA++L ++ ++K KL +++I++ + N DNDIKGISLSEFHVIECIGKNNMPNNIFIAKELNMTKGGISKINSKLLLKDIIRADKIEN 117
Sbict 58
                DERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL
Query 88
               D+R + +SLTEK + + K H HEK ++ + + EE I KFL L
DKREIYYSLTEKGIALFKLHEHLHEKEREKLMKILSNYKOEEITTILKFLEDL 170
Sbict 118
>ref|YP 808725.1| 🖪 transcriptional regulator [Lactococcus lactis subsp. cremoris
SK11]
 gb|ABJ72303.1| G Transcriptional regulator [Lactococcus lactis subsp. cremoris
SK111
Length=292
 GENE ID: 4432217 LACR 0742 | transcriptional regulator
[Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 51.2 bits (121), Expect = 2e-05, Method: Compositional matrix adjust. Identities = 35/91 (3%), Positives = 50/91 (54%), Gaps = 3/91 (3%)
               TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHH 111
TNA+IAE L I P++VT +K+L+E E++ + ND+RV LTEK + T H
TNAEIAELLDIKPSSVTAQVKQLEEAEMVIRKQDENDKRVNRIFLTEKGREAQETRDTMH 116
Query 52
Sbict 57
Query 112 EKTLSTYQELGNKFTDEEQEVISKFLSALTE 142 T GN TDEEQ+ ++ + L E
               NDISETI--FGN-LTDEEQKQLANLMEKLVE 144
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17
>ref|YP 001698248.1|  MarR family transcriptional regulator [Lysinibacillus sphaericus
C3-41]
  gb|ACA40118.1| C transcriptional regulator, MarR family (Lysinibacillus sphaericus
C3-41]
Length=162
GENE ID: 6022244 Bsph 2567 | MarR family transcriptional regulator
[Lysinibacillus sphaerTcus C3-41] (10 or fewer PubMed links)
 Score = 51.2 bits (121), Expect = 3e-05, Method: Compositional matrix adjust. Identities = 36/111 (32%), Positives = 60/111 (54%), Gaps = 1/111 (0%)
                Ouerv 36
Sbjct 44
                SLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ 145
LT ++ H HEK + Y L + F +E E I FL +T++ +
LLTAFGKQLSSIHEQLHEKARNRYLRLLDSFNTDELETIVTFLEMITDKLK 154
Query 95
Sbjct 104
>ref|NP_266864.1| 🖸 trans<u>cr</u>iption regulator [Lactococcus lactis subsp. lactis Il1403]
 gb|AAK04806.1|AE006304 6 G transcriptional regulator [Lactococcus lactis subsp. lactis Il1403]
Length=291
GENE ID: 1114333 rmaB | transcription regulator
[Lactococcus lactis subsp. lactis I11403] (10 or fewer PubMed links)
 Score = 51.2 bits (121), Expect = 3e-05, Method: Compositional matrix adjust. Identities = 35/91 (3%), Positives = 50/91 (54%), Gaps = 3/91 (3%)
                Sbict 57
                EKTLSTYQELGNKFTDEEQEVISKFLSALTE
Query 112
                T GN TDEEQE ++ + L E
NDISETI--FGN-LTDEEQEQLAFLMEKLVE
Sbict 117
>ref[ZP_01723095.1| transcriptional regulator, MarR family protein [Bacillus sp.
gb|EAZ86562.1| transcriptional regulator, MarR family protein [Bacillus sp. B14905]
B14905]
Length=162
 Score = 50.4 bits (119), Expect = 4e-05, Method: Compositional matrix adjust. Identities = 35/111 (31%), Positives = 60/111 (54%), Gaps = 1/111 (0%)
                TSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLW
T TQ HI+ + Q S N ++E L +S A+TKA+KK+ E+ +I +R +++ + +
Ouerv 36
                TLTQLHIVSAIKAQGSANNTFLSETLNVSKPAITKAIKKMLEKNVIVETRQEANQKEIHY
Sbjct 44
               SLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ 145
LT ++ H HEK + Y L + F +E E I FL +T++ +
LLTAFGKQLSSIHEQLHEKARNRYLRLLDSFNTDELETIITFLEMITDKLK 154
Query 95
Sbjct 104
>ref|YP 808147.1|  transcriptional regulator [Lactococcus lactis subsp. cremoris
 gb|ABJ71725.1| G Transcriptional regulator [Lactococcus lactis subsp. cremoris
SK111
Length=169
GENE ID: 4432499 LACR 0099 | transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 50.4 bits (119), Expect = 5e-05, Method: Compositional matrix adjust. Identities = 27/98 (27%), Positives = 55/98 (56%), Gaps = 0/98 (0%)
                HILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA 100
Query 41
                HIL L ++ T ++A KL ++ VT+A++ L + + + + +A ND++ + + +T K
HILSALTKKDLTGIELATKLSVTRGGVTRAVQNLIKHQFLTTYQADNDKKKIYYHITTKG 112
Sbjct 53
Query 101
                VPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS
               + VA H H+ ++ +K+ + E+ +I FLS
LKVASIHDKMHKIMDLKLGQIFDKYNENEKSIILNFLS
>ref[NP_786298.1]  transcription regulator [Lactobacillus plantarum WCFS1]
emb|CAD65154.1| G transcription regulator [Lactobacillus plantarum WCFS1]
Length=178
GENE ID: 1063436 lp 2967 | transcription regulator [Lactobacillus plantarum WCFS1] (10 or fewer PubMed links)
 Score = 50.1 bits (118), Expect = 6e-05, Method: Compositional matrix adjust. Identities = 33/104 (31%), Positives = 57/104 (54%), Gaps = 5/104 (4%)
               QEHILMLLAEQIS--TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSL 96 Q IL +LA+ + TNA+IAE L I P++V+ L +L++ LI+ + +D+RVV+ L QMGILRVLADAPAGLTNAEIAEILDIRPSSVSATLNRLEDGGLIEREPSAHDKRVVIVRL 102
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Query 39 Sbjct 43

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Ouerv 97
              TEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL 140
              +++ +A A + T +L TD+E+ + L L
SDRGREMADHRA---QGTSDLADQLFGNLTDDERNQLQHLLDKL 143
>ref|YP_757477.1|  MarR family transcriptional regulator [Maricaulis maris MCS10]
 gb|ABI66539.1| G transcriptional regulator, MarR family [Maricaulis maris MCS10]
 GENE ID: 4285355 Mmar10_2247 | MarR family transcriptional regulator
[Maricaulis maris MCS10]
 Score = 50.1 bits (118), Expect = 6e-05, Method: Compositional matrix adjust. Identities = 39/146 (26%), Positives = 75/146 (51%), Gaps = 5/146 (3%)
              MSLANQIDQFLGTIMQFAENK-HEILLGKCESDVKLTSTQEHILMLLAEQISTN-AKIAE
Querv 1
              M+ A +D+ L +++ A K + + +S +TS Q +L LLA + I E MARARAVDRRLFLLLEIAARKLNRDADARLKSVAGVTSAQAAVLFLLARRGERRMGDIGE
Sbjct 1
              Query 59
Sbict 61
Query 119
              QELGNKFTDEEQEVISKFLSALTEEF 144
                       +E+ +++ + L+ +
              -ELENRLGEEDADMLHRVLTRIAVDF 143
Sbjct 119
>ref|NP_391385.1| 🖸 hypothetical protein BSU35050 [Bacillus subtilis subsp. subtilis
str. 168)
ref[ZP_03593302.1] hypothetical protein Bsubsl_18971 [Bacillus subtilis subsp. subtilis
str. 168]
ref[ZP 03597587.1| hypothetical protein BsubsN3_18887 [Bacillus subtilis subsp. subtilis str. NCIB 3610]
ref[ZP 03601991.1| hypothetical protein BsubsJ_18850 [Bacillus subtilis subsp. subtilis
 ref|ZP_03606276.1| hypothetical protein BsubsS 19006 [Bacillus subtilis subsp. subtilis
str. SMY]
sp|034692.1|YVNA_BACSU RecName: Full=Uncharacterized HTH-type transcriptional regulator
 gb|AAC67281.1| YvnA [Bacillus subtilis]
 GENE ID: 936639 vvnA | vvnA [Bacillus subtilis subsp. subtilis str. 168]
(10 or fewer PubMed links)
 Score = 50.1 bits (118), Expect = 6e-05, Method: Compositional matrix adjust. Identities = 30/104 (28%), Positives = 59/104 (56%), Gaps = 1/104 (0%)
              TSTQEHILMLL-AEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLW
Ouerv 36
              T TQ HI+ + Q N+ +A +L IS AAV+KA+ L + +I ++ +++ + + TLTQLHIVSCIHTSQNVNNSFLASRLHISKAAVSKAVHALLKHNIITVTKKPGNKKEIFY 107
Sbjct 48
              SLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS 138
             +LT+ +A H HEK Y++L N+F+ ++ + + F +
TLTDSGRKLAALHEQLHEKAKEQYKQLFNEFSIDDLKTVTAFFN 151
>ref|NP_111681.1|  transcription regulator (SlyA-related) [Thermoplasma volcanium
GSS1]
 Length=143
GENE ID: 1441302 TVN1162 | transcription regulator (SlvA-related)
[Thermoplasma volcanium GSS1] (10 or fewer PubMed links)
 Score = 49.7 bits (117), Expect = 7e-05, Method: Compositional matrix adjust. Identities = 30/81 (37%), Positives = 45/81 (55%), Gaps = 0/81 (0%)
Query 26
              LGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRA
             +G+ S + + IL LL+E ST K+AE ++PA +T L +++ Q LI SR+ MGESLSHISAKPIEVRILYLLSEDESTVNKLAELTDVTPAWITGTLDEMESQGLIVRSRS 86
Sbjct 27
Query 86
              TNDERVVLWSLTEKAVPVAKE
                D RVV
                          +TEK + V
              GEDRRVVNVHITEKGIEVLNE 107
Sbjct 87
>ref|NP_419215.1| G MarR family transcriptional regulator [Caulobacter crescentus
CB151
 ref|YP_002515775.1|  transcriptional regulator, MarR family [Caulobacter crescentus
NA10001
 gb|AAK22383.1| 😉 transcriptional regulator, MarR family (Caulobacter crescentus
CB15)
gb|ACL93867.1|  transcriptional regulator, MarR family [Caulobacter crescentus NA1000]
Length=147
GENE ID: 942105 CC_0396 | MarR family transcriptional regulator [Caulobacter crescentus CB15] (10 or fewer PubMed links)
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Score = 49.3 bits (116), Expect = 9e-05, Method: Compositional matrix adjust. Identities = 32/119 (26%), Positives = 63/119 (52%), Gaps = 4/119 (3%)
                   GKCESDVKLTSTQEHILMLLAEQIST-NAKIAEKLKISPAAVTKALKKLQEQELIKSSRA 85
Query 27
                  G+ ++ LT+ Q +L L E+ + A+ L ++P+A+T + ++ EL++
GRMAAEGGLTAAQSGVLFFLGERDGALIGEAADALDLAPSAMTGLIDRMARAELVERRAD 85
Sbict 26
                   TNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF 144
Query 86
                  DR + LT+K A++ A + ++ +L FTDEE V+S++L++L +F
AKDGRAMHLHLTDKGR-AARDTAKAGLRGVNA--QLTEGFTDEEISVVSRWLASLQTKF 141
Sbjct 86
>ref|YP 001031482.1| C MarR family transcriptional regulator [Lactococcus lactis subsp.
cremoris MG1363]
 emb|CAL96727.1|  transcriptional regulator, MarR family [Lactococcus lactis subsp.
cremoris MG1363]
Length=172
GENE ID: 4798609 rmaD | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 49.3 bits (116), Expect = 1e-04, Method: Compositional matrix adjust. Identities = 26/98 (26%), Positives = 55/98 (56%), Gaps = 0/98 (0%)
                   HILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA 100
                  HIL L ++ T ++A KL ++ VT+A++ L + + + + + A ND++ + + + + T K
HILSALTKKDLTGIELATKLSVTRGGVTRAVQNLIKHQFLTTYQADNDKKKIYYHITTKG 115
Sbict 56
                  VPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS
Query 101
                 + VA H H+ ++ +K+ + +FLS
LKVASIHDKMHKIMDLKLGOIFDKYNENDKSIILNFLS
Sbict 116
>ref[NP 603578.1]  Mark family transcriptional regulator [Fusobacterium nucleatum subsp. nucleatum ATCC 25586]

gb[AAL94877.1]  Transcriptional regulator, Mark family [Fusobacterium nucleatum
subsp. nucleatum ATCC 25586)
Length=225
GENE ID: 991648 FN0681 | Mark family transcriptional regulator [Fusobacterium nucleatum subsp. nucleatum ATCC 25586] (10 or fewer PubMed links)
 Score = 48.9 bits (115), Expect = 1e-04, Method: Compositional matrix adjust. Identities = 29/88 (32%), Positives = 50/88 (56%), Gaps = 3/88 (3%)
                  Query 35
Sbjct 34
                  LWSLTEKAVPVAKEHATHHEKTLSTYQE 120
Ouerv 93
                  FVSLTKKGVDALTYHNNYHKMIMASITE 120
Sbjct 93
>ref|YP 138915.1|  MarR family transcriptional regulator [Streptococcus thermophilus
LMG 183T1]
 gb|AAV60100.1| 🖸 transcriptional regulator, MarR family [Streptococcus thermophilus
LMG 18311]
Length=144
GENE ID: 3164787 stu0381 | MarR family transcriptional regulator [Streptococcus thermophilus LMG 18311] (10 or fewer PubMed links)
 Score = 48.9 bits (115), Expect = 1e-04, Method: Compositional matrix adjust. Identities = 31/90 (34%), Positives = 51/90 (56%), Gaps = 3/90 (3%)
                  STNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATH 110
+T + +A +L ++ VT +L KL+++ I +R++ D RVV SL++K V + H
TTPSAVARELMLTLGTVTTSLNKLEKKGYIIRTRSSVDRRVVHLSLSKKGRLVYRLHRGF 109
Query 51
Sbjct 50
                  HEKTLSTYQELGNKFTDEEQEVISKFLSAL 140
H+ + T E F DEE +V+SK L L
HKSMVKTITE---GFNDEELKVMSKGLENL 136
Query 111
Sbjct 110
>ref|YP_002165566.1|  possible MarR family transcriptional regulator [Fusobacterium nucleatum subsp. polymorphum ATCC 10953]
gb|EDK89119.1|  possible MarR family transcriptional regulator [Fusobacterium nucleatum subsp. polymorphum ATCC 10953]
Length=225
GENE ID: 6818656 FNP 1336 | possible MarR family transcriptional regulator [Fusobacterium nuclea\overline{t}um subsp. polymorphum ATCC 10953]
 Score = 48.9 bits (115), Expect = 1e-04, Method: Compositional matrix adjust. Identities = 29/88 (32%), Positives = 50/88 (56%), Gaps = 3/88 (3%)
                  LTSTQEHILMLLAE--QISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVV 92
LT T+ HI+ + E Q++ N ++A+K+ I+ T A+ KL ++ I +R+T D R V
LTHTELHIIESIGENTQLTMN-ELADKIGITMGTATVAISKLSDKGYIDRARSTTDRKV 92
Ouerv 35
Sbjct 34
Query 93
                  LWSLTEKAVPVAKEHATHHEKTLSTYQE
                     SLT+K V
                                       H +H+ +++
```

```
>ref|ZP 00144611.1| Transcriptional regulator, MarR family [Fusobacterium nucleatum subsp. Vincentii ATCC 49256]
gb|EAA23791.1| Transcriptional regulator, MarR family [Fusobacterium nucleatum subsp. vincentii ATCC 49256]
Length=225
 Score = 48.9 bits (115), Expect = 1e-04, Method: Compositional matrix adjust. Identities = 29/88 (32%), Positives = 50/88 (56%), Gaps = 3/88 (3%)
               Sbjct 34
Query 93
                LWSLTEKAVPVAKEHATHHEKTLSTYQE 120
               SLT+K V H +H+ +++ E
FVSLTKKGVDALTYHNNYHKMIMASITE
Sbict 93
>ref|YP_819912.1| 🖸 MarR family transcriptional regulator [Streptococcus thermophilus
 gb|ABJ65716.1| ctranscriptional regulator, MarR family [Streptococcus thermophilus
Length=144
GENE ID: 4437359 STER 0428 | Mark family transcriptional regulator [Streptococcus thermophilus LMD-9] (10 or fewer PubMed links)
 Score = 48.9 bits (115), Expect = 1e-04, Method: Compositional matrix adjust. Identities = 31/90 (34%), Positives = 51/90 (56%), Gaps = 3/90 (3%)
               STNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATH 110
+T + +A +L ++ VT +L KL+++ I +R++ D RVV SL++K V + H
TTPSAVARELMLTLGTVTTSLNKLEKKGYIIRTRSSVDRRVVHLSLSKKGRLVYRLHRAF 109
Query 51
Sbict 50
               HEKTLSTYQELGNKFTDEEQEVISKFLSAL 140
Query 111
               H+ + T E F DEE +V+SK L L
HKSMVKTITE---GFNDEELKVMSKGLENL 136
Sbjct 110
>ref|YP_079349.1|  transcriptional regulator YvmB [Bacillus licheniformis ATCC 14580]
 gb|AAU23711.1|  possible transcriptional regulator YvmB (Bacillus licheniformis
ATCC 14580]
Length=161
GENE ID: 3031484 yvmB | transcriptional regulator YvmB [Bacillus licheniformis ATCC 14580] (10 or fewer PubMed links)
 Score = 48.5 bits (114), Expect = 2e-04, Method: Compositional matrix adjust. Identities = 35/117 (29%), Positives = 59/117 (50%), Gaps = 4/117 (3%)
               CESDVKL---TSTQEHILMLLA-EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSR 84
Query 29
               ESDVK T H++ + ++ N IA+K+ +S A +TK KL ++ LIK + AESDVKRLPGNMTTIHVISCIGHDEPINNTGIAKKMNLSKANITKISSKLLKEGLIKRFQ
Sbict 36
Ouerv 85
               Sbjct 96
               LTDNKKEIYFRLTPSGKQVFELHEKLHQQKADQFSRFLDSFSTAEQGAILKFLQGLT
>ref|ZP_01219563.1| putative transcriptional regulator, MarR family protein [Photobacterium
profundum 3TCK]
gb[EAS43796.1]
                     putative transcriptional regulator, MarR family protein [Photobacterium
profundum 3TCK)
Length=138
 Score = 48.1 bits (113), Expect = 2e-04, Method: Compositional matrix adjust. Identities = 39/143 (27%), Positives = 73/143 (51%), Gaps = 13/143 (9%)
Query 7
               IDQFLGTIMQFAEN----KHEILLGKCESDVKLTSTQEHILMLLAEQISTNAK-IAEKLK 61
               Sbjct 4
Query 62
               ISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQEL
                                                                                                 121
               ++ +T + K+ + \to LI+ +D R +L LTEK V + +\to H LS Q++ VTTGTLTVQVDKMVQAELIQRRPHESDRRSILVDLTEKGVEMYQEHDHLH---LSLTQDI
Sbjct 59
               GNKFTDEEQEVISKFLSALTEEF
Ouerv 122
               + D E++ + +L+ + +EF
TAQLDDVERKNLLMYLTKMNQEF 138
Sbjct 116
>ref|YP_091766.1|  YvmB [Bacillus licheniformis ATCC 14580]
 gb|AAU41073.1| G YvmB [Bacillus licheniformis DSM 13]
Length=163
GENE ID: 3097946 vvmB | YvmB [Bacillus licheniformis ATCC 14580] (10 or fewer PubMed links)
 Score = 48.1 bits (113), Expect = 2e-04, Method: Compositional matrix adjust. Identities = 35/117 (29%), Positives = 59/117 (50%), Gaps = 4/117 (3%)
```

21 Annex 1 Ouerv 29 CESDVKL---TSTQEHILMLLA-EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSR 84 ESDVK T H++ + ++ N IA+K++S A +TK KL ++ LIK +
AESDVKRLPGNMTTIHVISCIGHDEPINNTGIAKKMNLSKANITKISSKLLKEGLIKRFQ 97 Sbjct 38 ATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALT 141 Ouerv 85 T++++ + LT V + H H++ + + F+ EQ I KFL LT LTDNKKEIYFRLTPSGKQVFELHEKLHQQKADQFSRFLDSFSTAEQGAILKFLQGLT 154 Sbjct 98 >ref|YP 129221.1| G MarR family transcriptional regulator [Photobacterium profundum emb|CAG19419.1| G putative transcriptional regulator, MarR family (Photobacterium profundum SS9] Length=138 GENE ID: 3122954 PBPRA1008 | MarR family transcriptional regulator [Photobacterium profundum SS9] (10 or fewer PubMed links) Score = 48.1 bits (113), Expect = 2e-04, Method: Compositional matrix adjust. Identities = 39/143 (27%), Positives = 73/143 (51%), Gaps = 13/143 (9%) Query 7 IDQFLGTIMQFAEN----KHEILLGKCESDVKLTSTQEHILMLLAEQISTNAK-IAEKLK 61 THO TI++FE + ++ GK + Q HI+ +L + K +A+K+
IEQLNHTIIEFYEKLSSWEQSVVRGKG----FSLPQIHIVEILGAHGAMRMKELADKIG Sbict 4 Query 62 ISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQEL ++ +T + K+ + ELI+ +D R +L LTE+ V + +EH H LS Q++ VTTGTLTVQVDKMVQAELIQRRPHESDRRSILVDLTEQGVEMYQEHDHLH---LSLTQDI Sbjct 59 Query 122 GNKFTDEEQEVISKFLSALTEEF 144 K D E++ + +L+ + +EF
TAKLDDTERKNLLMYLTKMNQEF 138 Sbict 116 >ref|ZP\_00990845.1| hypothetical transcriptional regulator, MarR family [Vibrio splendidus gb|EAP94213.1| hypothetical transcriptional regulator, MarR family [Vibrio splendidus 12801] Length=301 Score = 48.1 bits (113), Expect = 2e-04, Method: Compositional matrix adjust. Identities = 35/117 (29%), Positives = 62/117 (52%), Gaps = 1/117 (0%) LLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSR 84 +L K D+ L Q H L+ L +Q T ++A+KL I + ++A+ L + LI++S MLDKDCGDIALPPIQAHTLIELEQQPLTVNQLADKLNIDKSNASRAVNNLAKNSLIOTSP Sbjct 19 Query 85 ATNDERVVLWSLTEKAV-PVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL 140 ND+R V+ S+TE+ + HPNDKRSVVASVTEQGIKTLAQLHSQQNQFYDSVLERLTEAETQQVSGGIKHYLKAL 135 Sbjct 79 >ref|YP\_001319891.1| 🖸 MarR family transcriptional regulator (Alkaliphilus metalliredigens gb|ABR48232.1| G transcriptional regulator, MarR family [Alkaliphilus metalliredigens OYMF1 Length=143 GENE ID: 5312143 Amet 2072 | MarR family transcriptional regulator [Alkaliphilus metalliredigens QYMF] Score = 47.8 bits (112), Expect = 3e-04, Method: Compositional matrix adjust. Identities = 27/67 (40%), Positives = 42/67 (62%), Gaps = 1/67 (1%) Query 35 LTSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL +T TQ L++L E+ AK + EKL + +T LKKL+ +ELI R+T DER+++
ITYTQYITLLVLWEKPKITAKELGEKLYLDSGTLTPLLKKLESKELITRKRSTKDERIMI 96 Sbict 37 Query 94 WSLTEKA 100 +LT+K VTLTDKG 103 Sbjct 97 gb|ABZ96926.1| Putative transcriptional regulator, MarR family [Leptospira biflexa serovar Patoc strain 'Patoc 1 (Paris)']
Length=133 GENE ID: 6222463 LEPBI\_I0798 | MarR family transcriptional regulator [Leotospira biflexa serovar Patoc strain 'Patoc 1 (Paris)'] (10 or fewer PubMed links) Score = 47.8 bits (112), Expect = 3e-04, Method: Compositional matrix adjust. Identities = 26/71 (36%), Positives = 45/71 (63%), Gaps = 1/71 (1%) SDVKLTSTQEHILMLLAEQ-ISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDE 89 +D+ LT Q +++++ E+ IST +KI +KL++ +T LK+L++ EL+ R NDE ADLGLTYPQYLVMLVMWEEKISTVSKIGDKLQLDSGTLTPLLKRLEQMELLDRMRNPNDE 85 Ouerv

Sbjct 26

Sbjct 86

Ouerv

90

RVVLWSLTEKA 100

RSVNIVLSKKG 96

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>ref|YP 108512.1| 🗲 putative transcriptional regulatory protein [Burkholderia pseudomallei
K96243)
 ref|YP 102754.1| G MarR family transcriptional regulator [Burkholderia mallei ATCC
23344]
 ref|ZP 00438156.1| COG1846: Transcriptional regulators [Burkholderia mallei GB8
horse 4T
 46 more sequence titles
ref|YP_333321.1| G MarR family transcriptional regulator [Burkholderia pseudomallei
 ref|YP_992831.1| G MarR family transcriptional regulator [Burkholderia mallei SAVP1]
 ref|YP_001026174.1| G MarR family transcriptional regulator [Burkholderia mallei NCTC
 ref|YP 001058785.1| G MarR family transcriptional regulator [Burkholderia pseudomallei
 ref|YP 001080556.1| G MarR family transcriptional regulator [Burkholderia mallei NCTC
102471
 ref|YP 001066038.1| 🖸 MarR family transcriptional regulator [Burkholderia pseudomallei
1106a1
                      transcriptional regulator, MarR family [Burkholderia pseudomallei
 ref|ZP 01769009.1|
 ref|ZP_02102319.1|
                       transcriptional regulator, MarR family protein [Burkholderia
pseudomallei 1106b)
ref|ZP_02109049.1
                       transcriptional regulator, MarR family protein (Burkholderia
pseudomallei 1710a]
 ref|ZP 02266811.1|
                      transcriptional regulator, MarR family protein [Burkholderia
mallei PRL-20]
ref[ZP_02403045.1]
                       transcriptional regulator, MarR family protein (Burkholderia
pseudomāllei DM98] ref[ZP 02411585.1|
                      transcriptional regulator, MarR family protein (Burkholderia
pseudomallei 14]
 ref|ZP 02447714.1|
                      transcriptional regulator, MarR family protein [Burkholderia
pseudomallei 91] ref[ZP 02455880.1]
                      transcriptional regulator, MarR family protein [Burkholderia
pseudomāllei 9)
 ref|ZP_02471448.1|
                       transcriptional regulator, MarR family protein [Burkholderia
pseudomallei B7210]
ref|ZP 02481920.1|
pseudomallei 7894]
                       transcriptional regulator, MarR family protein [Burkholderia
 ref|ZP_02490126.1|
                       transcriptional regulator, MarR family protein [Burkholderia
pseudomallei NCTC 13177)
ref|ZP 02498250.1|
pseudomallei 112|
                      transcriptional regulator, MarR family protein [Burkholderia
 ref|ZP 02506273.1|
                      transcriptional regulator, MarR family protein [Burkholderia
pseudomallei BCC215]
 ref|YP 002009903.1| C transcriptional regulator, MarR family [Burkholderia mallei ATCC
103991
 ref|YP_002036065.1|  transcriptional regulator, MarR family [Burkholderia pseudomallei
 ref|YP_002022669.1|  transcriptional regulator, MarR family [Burkholderia pseudomallei
Pasteur 52237]
 ref|YP 002050288.1| 🖸 transcriptional regulator, MarR family [Burkholderia pseudomallei
 ref|YP_002060819.1| G transcriptional regulator, MarR family [Burkholderia mallei FMH]
 ref[YP_002064980.1] G transcriptional regulator, MarR family [Burkholderia mallei JHU] ref[YP_002106328.1] G transcriptional regulator, MarR family [Burkholderia pseudomallei
 ref|YP 002110157.1| C transcriptional regulator, MarR family [Burkholderia mallei 2002721280] ref|ZP 03454874.1| transcriptional regulator, MarR family [Burkholderia pseudomallei
576]
 emb|CAH35912.1| G putative transcriptional regulatory protein (Burkholderia pseudomallei
K96243]
gb|AAU48847.1|  transcriptional regulator, MarR family [Burkholderia mallei ATCC 23344]
gb|ABA50381.1|  transcriptional regulator, MarR family (Burkholderia pseudomallei 1710b)
 gb|ABM52457.1| ctranscriptional regulator, MarR family [Burkholderia mallei SAVP1]
gb|ABN02741.1|  transcriptional regulator, MarR family [Burkholderia mallei NCTC 10229]
gb|ABN82036.1|  transcriptional regulator, MarR family [Burkholderia pseudomallei 668]
gb|ABN89050.1|  transcriptional regulator, MarR family [Burkholderia pseudomallei 1106a]
gb|ABO05799.1|  transcriptional regulator, MarR family (Burkholderia mallei NCTC 10247)
gb|EBA46431.1| transcriptional regulator, MarR family [Burkholderia pseudomallei 305]
 gb|EDK56373.1| 🖸 transcriptional regulator, MarR family (Burkholderia mallei FMH)
 gb|EDK60534.1|  transcriptional regulator, MarR family [Burkholderia mallei JHU]
 gb|EDK85546.1|  transcriptional regulator, MarR family [Burkholderia mallei 2002721280]
 gb|ED084870.1| 🖸 transcriptional regulator, MarR family (Burkholderia pseudomallei
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Annex 1 gb|EDO91962.1| G transcriptional regulator, MarR family (Burkholderia pseudomallei Pasteur 522371 gb|EEC33252.1| transcriptional regulator, MarR family [Burkholderia pseudomallei
576] Length=165 GENE ID: 3092472 BPSL1912 | putative transcriptional regulatory protein [Burkholderia pseudomallei K96243] (10 or fewer PubMed links) Score = 47.4 bits (111), Expect = 4e-04, Method: Compositional matrix adjust. Identities = 33/126 (26%), Positives = 69/126 (54%), Gaps = 5/126 (3%) Query 6 QIDQFLGTIMQFAEN-KHEILLGKCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKIS QI+ +G +M ++ ++ ++++++ TTQ +L +L +LA + ST A++A + I QINDSVGYLMSRVKSLMTNMVTQRTQTELGITGTQATMLFMLAVGKCSTAAELAREYGID Sbjct 17 PAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGN Query 64 +A+T+L +++++ L++ R++ D RVV LT++ + K L E+ + ASAITRLLDRVEKRGLLQRVRSSEDRRVVRLELTDEGRALTKRMPAIFRSVL---DEVLDSbjct 77 Query 124 KFTDEE 129 FT EE GFTPEE 139 Sbjct 134 >ref|YP 001961880.1| Transcriptional regulator, marR family [Leptospira biflexa serovar Patoc strain 'Patoc 1 (Ames)'] gb|ABZ93302.1| Transcriptional regulator, marR family [Leptospira biflexa serovar Patoc strain 'Patoc 1 (Ames)']
Length=145 GENE ID: 6388419 LBF\_0770 | Transcriptional regulator, marR family [Leptospira biflexa serovar Patoc strain 'Patoc 1 (Ames)'] (10 or fewer PubMed links) Score = 47.4 bits (111), Expect = 4e-04, Method: Compositional matrix adjust. Identities = 26/71 (36%), Positives = 45/71 (63%), Gaps = 1/71 (1%) SDVKLTSTQEHILMLL-AEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDE +D+ LT Q +++++ E+IST +KI +KL++ +T LK+L++ EL+ R NDE ADLGLTYPQYLVMLVMWEEKISTVSKIGDKLQLDSGTLTPLLKRLEQMELLDRMRNPNDE Ouerv 31 Sbict 38 Query 90 RVVLWSLTEKA 100 R V L++K RSVNIVLSKKG 108 Sbict 98 >ref|YP 001310724.1| G MarR family transcriptional regulator [Clostridium beijerinckii NCIMB 8052] gb|ABR35768.1| transcriptional regulator, MarR family [Clostridium beijerinckii NCIMB 8052] Length=155 **GENE ID: 5294814 Cbei 3649** | MarR family transcriptional regulator [Clostridium beijerinc $\overline{k}$ ii NCIMB 8052] Score = 47.4 bits (111), Expect = 4e-04, Method: Compositional matrix adjust. Identities = 30/109 (27%), Positives = 57/109 (52%), Gaps = 1/109 (0%) Query 34 KLTSTQEHILMLLAEQISTN-AKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVV KLT +Q H + + N K++++L ++ A+TK KKL + + + + + V
KLTFSQIHCIAAIEYIEDANITKLSQELGMTTGAITKMCKKLLNEGYVSKYOKEGNNKEV 87 Sbjct 28 Query 93 LWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALT 141 + LTE + V + H HEK+ + +++ ++ DEE+ I KFL +
YYDLTELGLNVCEIHNRIHEKSYNKKKDIIAQYNDEEKATILKFLHDMN 136 Sbjct 88 >ref|ZP\_01065868.1| hypothetical transcriptional regulator, MarR family protein [Vibrio sp. MED722]

gb|EAQ52745.1| hypothetical transcriptional regulator, MarR family protein [Vibrio sp. MED222] Length=301 Score = 47.0 bits (110), Expect = 5e-04, Method: Compositional matrix adjust. Identities = 26/77 (33%), Positives = 45/77 (58%), Gaps = 0/77 (0%) LLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSR 84 Query 25 +L K D+ L Q H L+ L +Q T ++A+KL I + ++A+ L + LI++S MLDKDCGDIALPPIQAHTLIELEQQPLTVNQLADKLNIDKSNASRAVNNLAKNSLIQTSP 78 Sbict 19 Query 85 ATNDERVVLWSLTEKAV ND+R V+ S+TE+

HPNDKRSVVASVTEOGI

Sbjct 79

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>ref|YP_140799.1| G MarR family transcriptional regulator [Streptococcus thermophilus
CNRZ1066]
  gb|AAV61984.1| 🖸 transcriptional regulator, MarR family [Streptococcus thermophilus
CNR21066)
Length=144
  GENE ID: 3167129 str0381 | MarR family transcriptional regulator
 [Streptococcus thermophilus CNRZ1066] (10 or fewer PubMed links)
  Score = 47.0 bits (110), Expect = 5e-04, Method: Compositional matrix adjust. Identities = 30/90 (33%), Positives = 50/90 (55%), Gaps = 3/90 (3%)
                 STNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATH 110
+T + +A +L ++ VT +L KL+++ I +R++ D RVV SL++K V + H
TTPSAVARELMLTLGTVTTSLNKLEKKGYIIRTRSSVDRRVVHLSLSKKGRLVYRLHRGF 109
Query 51
Sbjct 50
                 HEKTLSTYQELGNKFTDEEQEVISKFLSAL 140
H+ S + F DEE +V+SK L L
HK---SMVMRITEGFNDEELKVMSKGLENL 136
Query 111
Sbjct 110
>ref|YP_143999.1| G MarR family transcriptional regulator [Thermus thermophilus HB8]
 dbj|BAD70556.1| G transcriptional regulator MarR family [Thermus thermophilus HB8]
GENE ID: 3168756 TTHA0733 | MarR family transcriptional regulator [Thermus thermophilus HB8]
 Score = 46.6 bits (109), Expect = 6e-04, Method: Compositional matrix adjust. Identities = 34/109 (31%), Positives = 55/109 (50%), Gaps = 5/109 (4%)
                 LTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLW 94
L+ + H+L LLA+ + +++AE L++ P+ V+ L L+E+ L+K S D R V
LSPRKAHLLGLLAKGVDLPSQLAELLEVHPSQVSHLLAALEEEGLVKRSPDPQDRRKVKL 95
Query 35
Sbjct 36
Query 95
                 SLTEKAVPVAKEHATHHEKT-LSTYQELGNKFTDEEQEVISKFLSALTE
                                                                                         142
                 LT P +E A E L+ + + + EEQ + L LTE
FLT---PKGREAAARTEALWLAVFGRRLARLSPEEQAAFLRILRKLTE 140
Sbjct 96
>ref|YP_293371.1|_  regulatory protein, MarR [Ralstonia eutropha JMP134]
 gb|AAZ65514.1| G regulatory protein, MarR [Ralstonia eutropha JMP134]
Length=150
GENE ID: 3607869 Reut C6203 | regulatory protein, MarR
[Ralstonia eutropha JMF134]
 Score = 46.6 bits (109), Expect = 7e-04, Method: Compositional matrix adjust. Identities = 32/112 (28%), Positives = 58/112 (51%), Gaps = 1/112 (0%)
Query 6
                 QIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNA-KIAEKLKISP
                 +I Q +G + A N + + D+ +T Q IL+ L + ++T ++++ L I
RITQSVGFFLNRARNTLLMEMDAALKDLDITGQQMGILLSLTQGVATTPFELSKVLGIDT 74
Sbjct 15
Query 65
                 AAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLS
                +T+ L KL+ + L+ SR+ +D RVV +LT+K VA+ K L+ GLMTRMLDKLETKGLLSRSRSLDDRRVVNLTLTQKGQEVAERAPVVAPKVLN
Sbjct 75
>ref[YP_001307422.1]    MarR family transcriptional regulator [Clostridium beijerinckii
NCIMB 80521
gb|ABR32466.1|  transcriptional regulator, MarR family [Clostridium beijerinckii NCIMB 8052]
Length=153
GENE ID: 5291510 Cbei 0278 | MarR family transcriptional regulator [Clostridium beijerinc\overline{\text{K}}ii NCIMB 8052]
 Score = 46.6 bits (109), Expect = 7e-04, Method: Compositional matrix adjust. Identities = 30/114 (26%), Positives = 62/114 (54%), Gaps = 1/114 (0%)
                DVKLTSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRATNDER
D+ L ++ H++ + + NA I+++L ++ A++K KL ++ELIK + N+++
DMGLMLSEIHVIDCIGKNQLINATFISKELNMTKGAISKITSKLLKKELIKGNHLENNKK
Ouerv 32
Sbjct 40
                 VVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF
Query 91
+ ++LT + V K H H+ + ++ +K+ EE +I+ FL L E
Sbjct 100 EIYYTLTAQGKEVFKVHEILHKIESEKFVKILSKYDKEELSIINSFLEDLISEL
>ref|ZP_02950515.1| transcriptional regulator, MarR family [Clostridium butyricum
5521] gb|EDT74473.1| transcriptional regulator, MarR family [Clostridium butyricum 5521]
Length=145
 Score = 46.2 bits (108), Expect = 8e-04, Method: Compositional matrix adjust. Identities = 32/86 (37%), Positives = 48/86 (55%), Gaps = 2/86 (2%)
Query 17
                FAENKHEILLGK-CESDVKLTSTQE-HILMLLAEQISTNAKIAEKLKISPAAVTKALKKL 74
                +A ++ I L K C LT TQ +L+L ++ ST +I ++L + +T LKK+
YAASREVIKLYKPCLDKFNLTYTQYVAMLVLWEDEKSTVKEIGKRLHLDSGTLTPLLKKM 78
Sbjct 19
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QEQELIKSSRATNDERVVLWSLTEKA 100
+ ELIK R ND+RVV+ L EK
ESMELIKRYRDINDDRVVIVELAEKG 104
Query 75
Sbjct 79
>ref[NP_266269.1|  transcription regulator [Lactococcus lactis subsp. lactis Il1403]
 gb|AAK04211.1|AE006249_8  transcriptional regulator [Lactococcus lactis subsp. lactis Il1403]
GENE ID: 1113719 rmaD | transcription regulator
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
  Score = 46.2 bits (108), Expect = 0.001, Method: Compositional matrix adjust. Identities = 26/98 (26%), Positives = 55/98 (56%), Gaps = 0/98 (0%)
                          \begin{array}{llllllllaeq | limitate | lim
Sbjct 53
Query 101
                         VPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS 138
                        VA H H+ ++ +K+ ++E+ +I FLS
RKVATIHDKMHKIMDIRLGOIFDKYNEOEKSIILSFLS 150
Sbict 113
>ref|YP_002352539.1|  transcriptional regulator, MarR family [Dictyoglomus turgidum
  gb|ACK41925.1| G transcriptional regulator, MarR family (Dictyoglomus turgidum
DSM 6724]
Length=145
GENE ID: 7081680 Dtur 0640 | transcriptional regulator, MarR family
[Dictyoglomus turgidum DSM 6724]
  Score = 46.2 bits (108), Expect = 0.001, Method: Compositional matrix adjust. Identities = 33/118 (27%), Positives = 63/118 (53%), Gaps = 8/118 (6%)
                         SDVKLTSTQEHILMLLAEQIS-TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDE 89
S +KL Q IL+LL+E+ T +I E +KI P+ V +++++++ L+ + R D+
SGLKLYRGQAPILLLLSERDGLTQKEIVENMKIKPSTVAIMIRRMKKRGLVITKRDEKDK 85
Query 31
Sbjct 26
Query 90
                         RVVLWSLTEKAVPVAKEHATHHEKTLSTYQE--LGNKFTDEEQEVISKFLSALTEEFQ
                         R LT++ ++ +KT +E GN FT+EE+E + +L + + + RFSKVYLTDE----GRKFICKLKKTYKQLEEECFGN-FTEEERETLKNYLERIRDNLR 138
Sbict 86
>ref|ZP_01854255.1| probable mark-family transcription regulator {Planctomyces maris
gb|EDL59963.1| probable marR-family transcription regulator [Planctomyces maris DSM 8797]
Length=159
  Score = 46.2 bits (108), Expect = 0.001, Method: Compositional matrix adjust. Identities = 29/83 (34%), Positives = 46/83 (55%), Gaps = 2/83 (2%)
                         LGKCESDVKLTSTQEHILMLLA--EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSS 83
Ouerv 26
                         G+ + LT +Q ++L +L + + + +IA ++ A+T L +LQ QELIK +
FGRLFREYGLTPSQYNVLRILRGERKPMPSLEIANRMIQVVPAITGLLDRLQAQELIKRN 97
Sbict 38
Query 84
                         RATNDERVVLWSLTEKAVPVAKE 106
                        R T D RVV +T KA+ + KE
RCTEDRRVVYIEITAKALKLLKE
Sbict 98
>ref|YP_773280.1| 🖸 MarR family transcriptional regulator [Burkholderia ambifaria
AMMD1
 gb|ABI86946.1| 🖸 transcriptional regulator, MarR family [Burkholderia ambifaria
AMMD]
Length=164
  GENE ID: 4309612 Bamb 1388 | MarR family transcriptional regulator
[Burkholderia ambifaria AMMD]
 Score = 45.8 bits (107), Expect = 0.001, Method: Compositional matrix adjust. Identities = 31/118 (26%), Positives = 59/118 (50%), Gaps = 4/118 (3%)
                        KCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRAT 86 + + + + + T TQ +L ++A + ST A++A + I +AVT+ L +++++ L+ R+
Query 28
                         RTQEELGITGTQASMLFMIAVGRCSTAAELAREYGIDASAVTRLLDRVEKRGLLSRVRSI
Sbjct 40
Query 87
                         NDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF
                        D RVV LT++ +A+ L EL FT EE + L + + EDRRVVRLELTDEGRALAERLPPVFRSVL---DELLGGFTPEEVGFLKSMLRRILSNY
Sbjct 100
>ref[ZP 01131766.1] transcriptional regulator, MarR family protein [Pseudoalteromonas
tunicata D2]
gb|EAR30132.1| transcriptional regulator, Mark family protein [Pseudoalteromonas
tunicata D2]
Length=163
 Score = 45.8 bits (107), Expect = 0.001, Method: Compositional matrix adjust. Identities = 29/83 (34%), Positives = 47/83 (56%), Gaps = 1/83 (1%)
```

KCESDVKLTSTQEHILMLLAEQISTNAK-IAEKLKISPAAVTKALKKLQEQELIKSSRAT 86

Query 28

```
Annex 1
               K + LTS Q +L +++Q K IAE++ +S A +T L +L+ +EL+ R+T KLSKETGLTSPQLLVLQAISQQDGVMVKEIAEQINLSSATITSILDRLEIRELVIRERST 83
Sbict 24
                NDERVVLWSLTEKAVPVAKEHAT 109
Query 87
               D+R V SLT+K + K+ T
TDKRRVGISLTDKGFDIIKDSPT 106
Sbjct 84
>ref(ZP 00238819.1| transcriptional regulator, MarR family [Bacillus cereus G9241]
gb(EALT3614.1| transcriptional regulator, MarR family [Bacillus cereus G9241]
Length=152
 Score = 45.8 bits (107), Expect = 0.001, Method: Compositional matrix adjust. Identities = 27/85 (31%), Positives = 49/85 (57%), Gaps = 1/85 (1%)
                LTSTQEHILMLLAEQISTNA-KIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93
               LT Q +1L +L ++ A K+A+K+ + P+A+T + +L +QEL++ +D RVV+
LTPPQFY1LK1LDHYGASRATKLAKKMYVKPSAITVMIDRLIDQELVERYHDKDDRRVVI 94
Sbjct 35
               WSLTEKAVPVAKEHATHHEKTLSTY 118
LT+K +E T + ++ Y
Query 94
Sbjct 95
                IELTKKGKARVEEAMTARNEHIAKY
>ref[YP_002467440.1]  transcriptional regulator, MarR family [Candidatus Methanosphaerula
palustrīs E1-9c]
 gb|ACL17717.1| G transcriptional regulator, MarR family [Candidatus Methanosphaerula
palustris E1-9c}
Length=159
 GENE ID: 7271351 Mpal_2438 | transcriptional regulator, MarR family
[Candidatus Methanosphaerula palustris E1-9c]
 Score = 45.8 bits (107), Expect = 0.001, Method: Compositional matrix adjust. Identities = 25/85 (29%), Positives = 47/85 (55%), Gaps = 0/85 (0%)
               Query 56
Sbjct 53
Query 116
               STYQELGNKFTDEEQEVISKFLSAL 140
Sbjct 113 TRTTERVGPLSEEELELIARFFNAF 137
>ref|ZP 03547491.1| hypothetical protein BLAHAN 01259 [Blautia hansenii DSM 20583]
gb|EED60137.1| hypothetical protein BLAHAN_01259 [Blautia hansenii DSM 20583]
Length=168
 Score = 45.4 bits (106), Expect = 0.001, Method: Compositional matrix adjust. Identities = 32/105 (30%), Positives = 53/105 (50%), Gaps = 4/105 (3%)
               HILMLLAEQISTN-AKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEK
Query 41
               HI+ + E+ + N + +A+ L ++ +T A+ L ++ + R+ D RVVL SLT K
HIMEAIGEENAKNMSSVAKLLSVTVGTLTIAINGLVKKGYVARERSEEDRRVVLISLTGK 119
Sbict 60
Query 100
               {\tt AVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF}
               + H H+ + Q L DE+QE++ K L L E F
GRKANEHHKKFHDGMI---QALLKDLDDEQQEILVKSLLNLREFF 161
Sbict 120
>ref|ZP 02891041.1| transcriptional regulator, MarR family [Burkholderia ambifaria
IOP40-10]
gb|EDT03378.1| transcriptional regulator, MarR family (Burkholderia ambifaria IOP40-10)
Length=163
 Score = 45.4 bits (106), Expect = 0.001, Method: Compositional matrix adjust. Identities = 31/118 (26%), Positives = 59/118 (50%), Gaps = 4/118 (3%)
               KCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRAT 86
Query 28
               + + + + + TTQ +L ++A + ST A++A + I +AVT+ L +++++ L+ R+
RTQEELGITGTQASMLFMIAVGKCSTAAELAREYGIDASAVTRLLDRVEKRGLLSRVRSI
Sbjct 40
               Query 87
Sbjct 100
               EDRRVVRLELTDEGRELAERLPPVFRSVL---DELLGGFTPEEVGFLKSMLRRILSNY
>ref|YP 001808132.1| 🖸 MarR family transcriptional regulator [Burkholderia ambifaria
MC40-61
gb|ACB63916.1|  transcriptional regulator, MarR family [Burkholderia ambifaria MC40-6]
Length=164
GENE ID: 6177160 BamMC406 1428 | MarR family transcriptional regulator [Burkholderia ambifaria MC\overline{4}0-6]
 Score = 45.4 bits (106), Expect = 0.001, Method: Compositional matrix adjust. Identities = 31/118 (26%), Positives = 59/118 (50%), Gaps = 4/118 (3%)
               KCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRAT 86
Query 28
               + + + + + T TQ +L ++A + ST A++A + I +AVT+ L +++++ L+ R+
RTQEELGITGTQASMLFMIAVGKCSTAAELAREYGIDASAVTRLLDRVEKRGLLSRVRSI 99
```

Sbjct 40

```
NDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF
                                                                                            144
Query 87
              D RVV LT++ +A+ L EL FT EE + L + + EDRRVVRLELTDEGRALAERLPPVFRSVL---DELLGGFTPEEVGFLKSMLRRILSNY
Sbjct 100
>ref|ZP 02906301.1| transcriptional regulator, MarR family [Burkholderia ambifaria
 gb|EDT42559.1| transcriptional regulator, MarR family [Burkholderia ambifaria
MEX-51
Length=164
 Score = 45.4 bits (106), Expect = 0.001, Method: Compositional matrix adjust. Identities = 31/118 (26%), Positives = 59/118 (50%), Gaps = 4/118 (3%)
               KCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRAT 86
Query 28
               + + ++ +T TQ +L ++A + ST A++A + I +AVT+ L +++++ L+ R+
RTQEELGITGTQASMLFMIAVGKCSTAAELAREYGIDASAVTRLLDRVEKRGLLSRVRSI 99
Sbjct 40
Query 87
              NDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF
D RVV LT++ +A+ L EL FT EE + L + +
EDRRVVRLELTDEGRALAERLPPVFRSVL---DELLGGFTPEEVGFLKSMLRRILSNY
Sbjct 100
>ref|ZP 02362996.1| transcriptional regulator, MarR family protein [Burkholderia
oklahomensis C6786]
Length=165
 Score = 45.4 bits (106), Expect = 0.001, Method: Compositional matrix adjust. Identities = 31/126 (24%), Positives = 69/126 (54%), Gaps = 5/126 (3%)
               QIDQFLGTIMQFAEN-KHEILLGKCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKIS 63
QI+ +G +M ++ ++ ++++++ T TQ +L +LA + ST A++A + I
QINDSVGYLMSRVKSLMTNMVTQRTQTELGITGTQATMLFMLAVGKCSTAAELAREYGID 76
Querv 6
Sbjct 17
               PAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGN
Query 64
                                                                                               123
                +A+T+ L +++++ L++
                                        R++ D RVV
               ASAITRLLDRVEKRGLLQRVRSSEDRRVVRLELTDEGRDLTRRMPAIFRSVL---DQVLD
Sbjct 77
                                                                                              133
               KFTDEE 129
Ouerv 124
Sbjct 134 GFTPEE 139
>ref[NP 977055.1] G MarR family transcriptional regulator [Bacillus cereus ATCC 10987]
gb|AAS39663.1|  transcriptional regulator, MarR family [Bacillus cereus ATCC 10987]
Length=152
GENE ID: 2747926 BCE 0730 | MarR family transcriptional regulator [Bacillus cereus ATCC 10987] (10 or fewer PubMed links)
 Score = 45.4 bits (106), Expect = 0.002, Method: Compositional matrix adjust. Identities = 27/85 (31%), Positives = 49/85 (57%), Gaps = 1/85 (1%)
Query 35
               LTSTQEHILMLLAEQISTNA-KIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93
               LT Q +IL +L ++ A K+A+K+ + P+A+T + +L +QEL++ +D RVV+
LTPPQFYILKILDHYGASRATKLAKKMYVKPSAITVMIDRLIDQELVERYHDKDDRRVVV 94
Sbjct 35
               WSLTEKAVPVAKEHATHHEKTLSTY 118
Query 94
               LT+K +E T + ++ Y
IELTKKGKARVEEAMTARNEHIAKY 119
Sbjct 95
>ref[NP_782475.1] G MarR family transcriptional regulator [Clostridium tetani E88]
 gb|AAO36412.1| G transcriptional regulator, mark family [Clostridium tetani E88]
GENE ID: 1058804 marR | MarR family transcriptional regulator [Clostridium tetani E88] (10 or fewer PubMed links)
 Score = 45.4 bits (106), Expect = 0.002, Method: Compositional matrix adjust. Identities = 25/91 (27%), Positives = 50/91 (54%), Gaps = 0/91 (0%)
               STNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATH 110
Query 51
               + +IA L I+ +AV+K +++LQ++ LI SS+ ++++ + +LT + + K+H
ANGTQIANYLNITRSAVSKIIRRLQKENLIISSQKPDNKKEIFYTLTNEGNNIFKQHKQA 109
Sbjct 50
               HEKTLSTYQELGNKFTDEEQEVISKFLSALT 141
Query 111
HEK + + E+E + KFL
Sbjct 110 HEKWEIRDTKFLKTISTNEKETVFKFLKKFN 140
>ref|ZP_03570809.1| transcriptional regulator, MarR family (Burkholderia multivorans
CGD2M]
 ref|ZP 03577696.1| transcriptional regulator, MarR family [Burkholderia multivorans
CGD2
 ref[ZP_03586132.1| transcriptional regulator, MarR family [Burkholderia multivorans
gb|EED99839.1| transcriptional regulator, MarR family [Burkholderia multivorans CGD1]
 gb|EEE07966.1| transcriptional regulator, MarR family (Burkholderia multivorans
CĞD2]
gb|EEE14096.1| transcriptional regulator, MarR family [Burkholderia multivorans CGD2M]
Length=164
```

```
Score = 45.1 bits (105), Expect = 0.002, Method: Compositional matrix adjust. Identities = 32/118 (27%), Positives = 60/118 (50%), Gaps = 4/118 (3%)
                KCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRAT 86 + + ++ +T TQ +L ++A + ST A++A + I +AVT+ L +++++ L+ R+ RTQEELGITGTQASMLFMIAVGKCSTAAELAREYAIDASAVTRLLDRVEKRGLLCRVRSV 99
 Ouerv 28
 Sbjct 40
                 {\tt NDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF}
 Query 87
                D RVV LT++ +A+ L Q LG FT EE + L + + EDRRVVRLELTDEGRALAERLPAIFRSVLD--OLLGG-FTPEEVGFLKSMLRRILSNY
 Sbjct 100
 >ref|YP 001119315.1| 🖸 MarR family transcriptional regulator [Burkholderia vietnamiensis
gb|ABO54480.1|  transcriptional regulator, MarR family [Burkholderia vietnamiensis G4]
 Length=163
  GENE ID: 4953186 Bcep1808_1472 | MarR family transcriptional regulator
 [Burkholderia vietnamiensis G4]
 Score = 45.1 bits (105), Expect = 0.002, Method: Compositional matrix adjust. Identities = 35/126 (27%), Positives = 66/126 (52%), Gaps = 5/126 (3%)
                QIDQFLGTIMQFAENKHEILLG-KCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKIS 63
QI+ +G +M ++ L+ + + ++ +T TQ +L ++A + ST A++A + I
QINDSVGYLMSRVKSVMTNLVTQRTQEELGITGTQASMLFMIAVGKCSTAAELAREYGID 76
Query 6
Sbjct 17
                 {\tt PAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGN}
Query 64
                                            R+ D RVV
                                                            LT++
                                                                       +A+
                 ASAVTRLLDRVEKRGLLSRVRSIEDRRVVRLELTDEGRALAERLPPVFRSVL---DELLE
Sbjct 77
Query 124
                KFTDEE 129
                GFTPEE 139
Sbjct 134
>ref|YP_002418276.1|  Histone acetyltransferase HPA2 and related acetyltransferases [Vibrio splendidus LGP32]  emb|CAV19992.1|  Histone acetyltransferase HPA2 and related acetyltransferases
[Vibrio splendidus LGP32]
Length=301
 GENE ID: 7162224 VS 2735 | Histone acetyltransferase HPA2 and related
acetyltransferases [Vibrio splendidus LGP32]
 Score = 45.1 bits (105), Expect = 0.002, Method: Compositional matrix adjust. Identities = 35/117 (29%), Positives = 60/117 (51%), Gaps = 1/117 (0%)
                Query 25
Sbjct 19
Query 85
                ATNDERVVLWSLTEKA-VPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSAL 140
                ND+R V+ S+T++ +AK H ++ S + L T + I +L AL HPNDKRSVVASVTDQGKNTLAKLHNQQNQFYDSVLEHLTEAETQOVSGGIEHYLKAL
Sbjct 79
>ref|ZP 02355848.1|
                           transcriptional regulator, MarR family protein [Burkholderia
oklahomensis EO147]
Length=165
 Score = 45.1 bits (105), Expect = 0.002, Method: Compositional matrix adjust. Identities = 31/126 (24%), Positives = 69/126 (54%), Gaps = 5/126 (3%)
                QIDQFLGTIMQFAEN-KHEILLGKCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKIS
QI+ +G +M ++ ++ +++++ TQ +L +LA + ST A++A + I
QINDSVGYLMSRVKSLMTNMVTQRTQTELGITGTQATMLFMLAVGKCSTAAELAREYGID
Query 6
Sbjct 17
Query 64
                PAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGN
                +A+T+ L +++++ L++ R++ D RVV LT++ + + L ++ + ASAITRLLDRVEKRGLLQRVRSSEDRRVVRLELTDEGRELTRRMPAIFRSVL---DQVLD
Sbjct 77
                KFTDEE 129
Query 124
Sbjct 134
                GFTPEE
>ref|YP_443075.1| 🖸 MarR family transcriptional regulator (Burkholderia thailandensis
E2641
 ref[ZP 02374954.1]
                             transcriptional regulator, MarR family protein (Burkholderia
thailandensis TXDOH] ref[ZP_02388872.1]
                            transcriptional regulator, MarR family protein [Burkholderia
thailandensis Bt4
 gb|ABC39239.1|  transcriptional regulator, MarR family [Burkholderia thailandensis
E2641
Length=165
GENE ID: 3849294 BTH I2558 | Mark family transcriptional regulator [Burkholderia thailandensis E264] (10 or fewer PubMed links)
 Score = 44.7 bits (104), Expect = 0.002, Method: Compositional matrix adjust. Identities = 33/126 (26%), Positives = 67/126 (53%), Gaps = 5/126 (3%)
```

```
Ouerv 6
              QIDQFLGTIMQFAEN-KHEILLGKCESDVKLTSTQEHILMLLA-EQISTNAKIAEKLKIS
              QI+ +G +M ++ ++ ++++++++ T TQ +L +LA + ST A++A + I
QINDSVGYLMSRVKSLMTNMVTQRTQTELGITGTQATMLFMLAVGKCSTAAELAREYGID
Sbjct 17
              PAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGN
Ouerv 64
              +A+T+ L +++++ L++ R++ D RVV LT++ + K L E
ASAITRLLDRVEKRGLLQRVRSSEDRRVVRLELTDEGRELTKRMPEIFRSVLDQVLE---
Sbjct 77
              KFTDEE 129
Query 124
               FT EE
Sbjct 134
              GFTPEE 139
>ref|ZP 02633998.1| transcriptional regulator, MarR family [Clostridium perfringens
E str. JGS1987]
ref|ZP_02636384.1| transcriptional regulator, MarR family [Clostridium perfringens
B str. ATCC 3626)
ref[ZP_02643088.1]
                        transcriptional regulator, MarR family [Clostridium perfringens
NCTC 8239]
7 more sequence titles
ref|ZP_02629009.2| transcriptional regulator, MarR family [Clostridium perfringens
C str._JGS1495]
 ref[ZP 02954000.1] transcriptional regulator, Mark family [Clostridium perfringens
gb[EDS81585.1]
C str 702
D str. JGS1721)
                    transcriptional regulator, MarR family [Clostridium perfringens
 gb|EDT13394.1|
                    transcriptional regulator, MarR family [Clostridium perfringens
E str. JGS1987]
gb|EDT23419.1|
                    transcriptional regulator, MarR family [Clostridium perfringens
B str. ATCC 3626]
 gb|EDT71039.1|
                    transcriptional regulator, MarR family [Clostridium perfringens
D str. JGS1721]
 gb|EDT77904.1|
                    transcriptional regulator, MarR family [Clostridium perfringens
NČTC 82391
Length=154
 Score = 44.7 bits (104), Expect = 0.002, Method: Compositional matrix adjust. Identities = 33/99 (33%), Positives = 50/99 (50%), Gaps = 5/99 (5%)
              EQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEH
              E+ T +++A LKI+ +T A+ KL ++ + R D RVV+ LTEK K H EKARTMSEVALDLKITVGTLTTAINKLIKKGYVNRRRIEEDRRVVMIELTEKGTLAYKVH
Sbict 49
Query 108
              ATHHEKTLS-TYQELGNKFTDEEQEVISKFLSALTEEFQ 145
                             +ELG
                                         E+EV+
              EKFHEEMIDHVLEELGVS----EEEVLISSLDKLDKFFQ 143
Sbjct 109
>ref|YP_001374721.1|  MarR family transcriptional regulator (Bacillus cereus subsp.
cytotoxis NVH 391-98]

gb|ABS21726.1|  transcriptional regulator, MarR family [Bacillus cereus subsp.
cytotoxis NVH 391-98]
Length=136
GENE ID: 5344927 Bcer98 1405 | MarR family transcriptional regulator [Bacillus cereus subsp. Cytotoxis NVH 391-98]
 Score = 44.7 bits (104), Expect = 0.002, Method: Compositional matrix adjust. Identities = 37/119 (31%), Positives = 62/119 (52%), Gaps = 4/119 (3%)
              KCESDVKLTSTQEHILMLLAEQISTN-AKIAEKLKISPAAVTKALKKLQEQELIKSSRAT +CE + L+ IL + + T +KIA +L +S ++ +K+L +++LI R
Query 28
              RCEYENNLSHQAIRILQITSREAETTISKIASELNLSHNTASEHVKRLIQKDLILKERNK
Sbjct 16
Query 87
              NDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEFQ 145
                       +LT K
                                            EK L
                                                       L ++ + EEO++I +
Sbjct 76
              KDERVVNLALTAKGKEALIKHTLLDEKKLKI---LESQLSKEEQQIIEQAFSILAKEAQ
>ref|YP 819461.1| 🖸 transcriptional regulator [Leuconostoc mesenteroides subsp. mesenteroides
ATCC 82931
 gb|ABJ63088.1| 🖸 Transcriptional regulator [Leuconostoc mesenteroides subsp. mesenteroides
ATCC 8293]
Length=146
 GENE ID: 4423731 LEUM_2018 | transcriptional regulator
[Leuconostoc mesenteroides subsp. mesenteroides ATCC 8293] (10 or fewer PubMed links)
 Score = 44.7 bits (104), Expect = 0.002, Method: Compositional matrix adjust. Identities = 30/84 (35%), Positives = 46/84 (54%), Gaps = 5/84 (5%)
              KLTSTQEHILMLL-AEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVV 92
Query 34
              K+ STQ H+LMLL
              K+ STQ H+LMLL Q TN ++A + +S A+TKA+K L + + +D+R KINSTQAHLLMLLKILQSQTNTELAAAMNLSKPAITKAIKNLMRYHYVIAVVDDSDKRST
Sbict 32
Query 93
              LWSLTEKAVPVA----KEHATHHE 112
              + LTE +A + HAT H+
HYLLTEDGEKLAQLHEQAHATMHD
Sbjct 92
>ref[ZP_02027443.1| hypothetical protein EUBVEN_02713 [Eubacterium ventriosum ATCC
27560]
```

 $\mbox{\sc gb}|\mbox{\sc EDBVEN}\sc 02713$  [Eubacterium ventriosum ATCC 27560] Length=151

Score = 44.7 bits (104), Expect = 0.003, Method: Compositional matrix adjust. Identities = 27/90 (30%), Positives = 53/90 (58%), Gaps = 3/90 (3%)

IAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTL 115 IA+K+KI+ ++T ++ L ++ ++ ++ D R+V +LTEK + K H HEK IAKKMKITVGSLTTSMNSLVKKHYVERNRSEEDRRIVNITLTEKGIKAYKHHEEFHEK-- 117

Sbjct 60

Query 116 STYQELGNKFTDEEQEVISKFLSALTEEFQ 145 Q ++ +++E +V+ K L+ L++ F Sbjct 118 -MSQAAISEMSEDEVKVLLKSLNNLSKFFH 146

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SERVICE ADMINISTRATIF ERIC ANDRIEU
DIRECTEUR ADMINISTRATIF ET FINANCIER ANNIE LECUYER
RESPONSABLE ADMINISTRATIVE PL

**DOCUMENTATION** & INFORMATIOUE

FABRICE SAUSSEREAU DEUST DROIT INFORMATIQUE ET SYSTEME D'INFORMATION

S.A. au Capital de 360 000,00 € 380 823 179 R.C.S. Paris

O CONSEIL EN PROPRIÉTÉ INDUSTRIELLE O MANDATAIRE EN BREVETS EUROPÉENS O MANDATAIRE AGRÉÉ PRÉS L'OHP-11 O DIRÉMÉTE DU CEIN O IRISH & BRITISH PATENT ATTORNEY

Paris, March 5, 2009

UNITED STATES OF AMERICA - PATENT APPLICATION No. 10/525,449

filed on: August 29, 2003

NATIONAL DE LA RECHERCHE name of: INSTITUT AGRONOMIQUE.

« Zinc-regulated prokaryotic expression cassettes. »

Invention: POOUET Isabelle, LLULL Daniel

Y/Ref.: 1169-034

O/Ref.: MJP/DG/md - F053900116/US/PCT

Dear Sir,

We refer to your letter of October 31, 2008 transmitting us the 1st Office Action issued in relation with the above-referred application.

We have discussed this case with the applicant, and beg you to find herein enclosed our comments and instructions concerning this application.

Since we believe that an interview with the Examiner will be necessary in order to discuss the objections and the possible amendments to the claims, you may request a third month extension of time.

Please acknowledge receipt of this letter by return.

In the meantime, we are,

Very truly yours,

WW

M.J. VIALLE-PRESLES

**B. ORES** 

Encl.:

- Memorandum, and

- Annexes (only by e-mail).



U.S Application 10/1525449

# MEMORANDUM OF INSTRUCTIONS FOR RESPONDING THE OFFICE ACTION OF OCTOBER 7, 2008

## 1) Claims objections:

We beg you to amend the claims as requested by the Examiner. Concerning claim 15, it differs from claim 12 in that the expression cassette does not comprise the sequence encoding the ZitR repressor (cf § [0025] to [0027] of the instant application), and can be used to obtain constitutive expression of a protein of interest in bacterial strains wherein the endogenous ZitR repressor is inactivated (cf. § [0053] of the instant application)

# 2) Rejection under 35 USC § 112.

We wish to point out that the definition "at least 80% identity with the *Lactococcus lactis* ZitR protein GENBANK AAK06214" does not encompass a broad variety of proteins.

The results of BLAST searchs using GENBANK AAK06214 as a query are enclosed therein. A first search was performed against all the protein sequences available in the nr database (all known protein sequences, including the complete sequences of various bacteria). The enclosed results (Annex 1) show that the only proteins having more than 80% identity with GENBANK AAK06214 (which is the ZitR protein of Lactococcus lactis subsp. lactis Il1403), are the ZitR proteins of Lactococcus lactis subsp. cremoris SK11 (89% identity) and the ZitR proteins of Lactococcus lactis subsp. cremoris MG1363 (88% identity).

A second search was performed against the whole genome sequences of Lactococcus lactis subsp. lactis Il1403, Lactococcus lactis subsp. cremoris SK11 and Lactococcus lactis subsp. cremoris MG1363. The enclosed results (Annex 2) show that the only lactococcal proteins having more than 80% identity with GENBANK AAK06214, are the ZitR proteins of Lactococcus lactis subsp. cremoris SK11 (89% identity) and the ZitR proteins of Lactococcus lactis subsp. cremoris MG1363 (88% identity).

Therefore, obtaining a nucleotide sequence encoding a protein having more than 80% identity with GENBANK AAK06214 would not have required require undue experimentation for one of skill in the art. He would have easily obtained it by routine screening of a lactococcal DNA library with a probe derived from the nucleotide sequence encoding GENBANK AAK06214 (or from nucleotides 357-794 of SEQ ID NO:2), and would inevitably have found the orthologs of GENBANK AAK06214.

We beg you to request an interview with the Examiner in order to discuss this issue of enablement. If necessary, you may amend paragraph b) of claim 12 so as to



specify that the polypeptide is a lactococcal polypeptide, and/or to indicate a higher percent of identity (85% rather than 80%) with GENBANK AAK06214.

# 3) Rejection under 35 USC § 102.

We beg you to insert in claims 12 and 15 the same disclaimer as in claim 18, which excludes the cassette and vector of POQUET et al.

# **BLAST Basic Local Alignment Search Tool**

**Eormatting options Download** 

# gb|AAK06214| (145 letters)

Results for: gb|AAK06214.1 zinc transport transcriptional regulator [Lactococcus lactis subsp. lactis l1403](145aa) Your BLAST job specified more than one input sequence. This box lets you choose which input sequence to show BLAST results for.

1

#### Query ID

gi|12725171|gb|AAK06214.1|AE006439\_11

#### Description

zinc transport transcriptional regulator [Lactococcus lactis subsp. lactis II1403]

#### Molecule type

amino acid

## Query Length

145

#### Database Name

3 databases

#### Description

Program

BLASTP 2.2.19+ / 注述公5

#### Reference

Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402.

Reference - compositional score matrix adjustment

Stephen F. Altschul, John C. Wootton, E. Michael Gertz, Richa Agarwala, Aleksandr Morgulis, Alejandro A. Schäffer, and Yi-Kuo Yu (2005) "Protein database searches using compositionally adjusted substitution matrices", FEBS J. 272:5101-5109.

Other reports: Search Summary [Taxonomy reports] [Distance tree of results]

## **Search Parameters**

Program	blastp	
Word size	3	
Expect value	10	
Hitlist size	100	
Gapcosts	11,1	
Matrix	BLOSUM62	
Threshold	11	
Composition-based stats 2		
Low Complexity Filter	Yes	
Filter string	L;	
Genetic Code	1	
Window Size	40	

#### **Database**

Posted date Mar 2, 2009 5:57 PM

Number of letters 2,055,736 Number of sequences 7,259 Entrez query none

## Karlin-Altschul statistics

Params	Ungapped	Gapped
Lambda	0.310848	0.267
K	0.124161	0.041

H 0.32399 0.14

## **Results Statistics**

Length adjustment 78 .

Effective length of query 67 .

Effective length of database 1489534 .

Effective search space 99798778 .

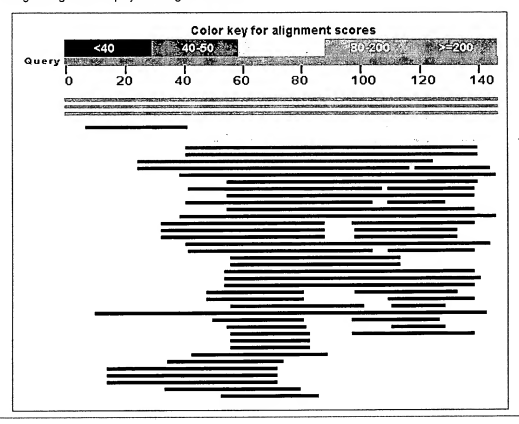
Effective search space used 99798778 .

Graphic Summary

### Distribution of 58 Blast Hits on the Query Sequence

[?]

An overview of the database sequences aligned to the query sequence is shown. The score of each alignment is indicated by one of five different colors, which divides the range of scores into five groups. Multiple alignments on the same database sequence are connected by a striped line. Mousing over a hit sequence causes the definition and score to be shown in the window at the top, clicking on a hit sequence takes the user to the associated alignments. New: This graphic is an overview of database sequences aligned to the query sequence. Alignments are color-coded by score, within one of five score ranges. Multiple alignments on the same database sequence are connected by a dashed line. Mousing over an alignment shows the alignment definition and score in the box at the top. Clicking an alignment displays the alignment detail.



Sequences producing significant alignments:	Score (Bits)	E Value
ref[NP 268273.1] zinc transport transcription regulator [Lact	291	2e-80 <b>G</b>
ref[YF S11979.]] transcriptional regulator [Lactococcus lacti	_68	2e-73 <b>G</b>
ref(YP 001033643.1) transcriptional regulator of the zit oper	267	3e-73 <b>G</b>
1ef[YP 001033:31.1] MarR family transcriptional regulator [La	<u> 53.3</u>	6e-09 <b>G</b>
ref[YP 808325.1] transcriptional regulator [Lactococcus lacti	<u> </u>	4e-08 <b>€</b>
refine 256364.1 transcription regulator [Lactococcus lactis	<u>51.2</u>	4e-08 <b>G</b>
<pre>:ef[YF 008147.1] transcriptional regulator [Lactococcus lacti</pre>	<u>50.4</u>	7e-08 <b>G</b>
<pre>ref[YP 001031462.1] MarR family transcriptional regulator [La</pre>	49.3	1e-07 <b>G</b>
ref[NP 2662(9.1] transcription regulator [Lactococcus lactis	35.	1e-06 <b>G</b>
ref[NP_067676.1] transcription regulator [Lactococcus lactis	41.3	4e-05 <b>G</b>
ref[7F 001032345,]] MarR family transcriptional regulator [La	<u>38.9</u>	2e-04 <b>G</b>
ref[YF 808479.1] transcriptional regulator [Lactococcus lacti	<u>37.0</u>	9e-04 <b>G</b>
mef[NP 066663.1] Mark family transcriptional regulator [Lacto	30.7	0.001 <b>G</b> 0.002 <b>G</b>
THE NE CONTACT. II transcription regulator [Lactococcus lactis	<u>35.8</u>	0.002 <b>G</b>
ref[YP 0003167" MarR family transcriptional regulator [La	35.5 25.5	0.002
ref[NP 00830] MarR family transcriptional regulator [Lacto ref[NP 008370] MarR family transcriptional regulator [Lacto	35,4	0.002 <b>G</b>
States and commensations and a state of the	27.3	0.003
AND ADDRESS OF THE PARTY OF THE	92.3	0.017
ref[YF 001031379.1] putative transcriptional regulator [Lacto ref[YF 008573.1] BadM/Rrf2 family transcriptional regulator [	32.3	0.018
ref[NP 266714.1] hypothetical protein L153086 [Lactococcus la	32.3	0.018 G
rof[YP 001033090.1] MarR family transcriptional regulator [La	31.6	0.031 <b>G</b>
ref[YP 808765.1] Mark family transcriptional regulator [Lacto	31.6	0.034 <b>G</b>
ref[YF 608795.2] MarR family transcriptional regulator [Lacto	31.2	0.044 <b>G</b>
ref[NF 165926.1] Mark family transcriptional regulator [Lacto	31.2	0.045 <b>G</b>
ref[YF 001031903.1] MarR family transcriptional regulator [La	28.9	0.18
ref[YF 001032520.1] MarR family transcriptional regulator [La	<u> 28.1</u>	0.31 <b>G</b>
ref[NF 167064.1] transcription regulator [Lactococcus lactis	23.1	0.35 <b>G</b>
<pre>ref[YF 001032533.1] transcriptional regulator (Lactococcus la</pre>	28.1	0.38 <b>G</b>
ref[YF 809312.1] Mn-dependent transcriptional regulator [Lact	23.1	0.38 <b>G</b>
ref[NF 267461.1] transcription regulator [Lactococcus lactis	<u> 27.7</u>	0.42 G
ref[NE 267696.1] transcription regulator [Lactococcus lactis	27.3	0.61 <b>G</b>
ref[NF 166753.1] NADPH-flavin oxidoreductase [Lactococcus lac	27.3	
ref[NF 267412.1] metalloregulator [Lactococcus lactis subsp	25.6	
ref[YF 001031697.1] putative cobalt ABC transporter ATP-bindi	26.2	1.2 G 2.4 G
ref[NF 166973.1] quinone oxidoreductase [Lactococcus lactis s ref[NF 168402.1] alkylphosphonate uptake protein [Lactococcus	<u>25.4</u> 25.4	2.4 <b>G</b>
the property of the state of th	24.6	3.4 <b>G</b>
ret[YF 001031533.1] transcriptional repressor CodY [Lactococcus	24.6	3.4 G
ret NF 266317.11 transcriptional repressor CodY (Lactococcus	24.6	3.5 G
ret NF 166838.11 intercellular adhesion protein [Lactococcus		3.6 <b>G</b>
rer[NF 266710.1] transcription regulator [Lactococcus lactis		3.6 G
rer[NF 166438.1] amino acid amidohydrolase [Lactococcus lacti	24.6	4.1 <b>G</b>
recure 809446. 1ysyl-tRNA synthetase [Lactococcus lactis su	24.6	4.3 <b>G</b>
ref[YP 001031741.1] lysyl-tRNA synthetase [Lactococcus lactis		4.7 <b>G</b>
rec[MF 266529.1] lysyl-tRNA synthetase [Lactococcus lactis su		4.7 <b>G</b>
reflyr 001031907.   NADPH-flavin oxidoreductase [Lactococcus		5.2 <b>G</b>
MADPH-flavin oxidoreductase [Lactococcus lac		<sub>5.5</sub> G <sub>6.1</sub> G
ref[M7 263/93.] exported serine protease [Lactococcus lactis		73
refire (01033660.1) housekeeping protease [Lactococcus lactis		
refire &11995.1  trypsin-like serine protease [Lactococcus la		
refine 269332.11 hypothetical protein L35545 [Lactococcus lac	25.5	7.5

		Annex 2
ref[NP 068337.1] hypothetical protein L39650 [Lactococcus lac	23.5	7.7 <u>G</u>
<pre>ref[YF 001032772.1] superfamily II DNA/RNA helicase [Lactococ</pre>	23.5	7.8 <b>G</b>
ref[YF 001033122.1] quinone oxidoreductase [Lactococcus lacti	23.5	9.3
ref(YF 000733.1) quinone oxidoreductase [Lactococcus lactis s	23.5	9.3 <b>G</b>
rof[Y5-795521.1] hypothetical protein LACR_C57 [Lactococcus 1	23.5	9.5 <b>G</b>
rgf[YP 812095.1] hypothetical protein LACR_2554 [Lactococcus	23.1	10.0 <b>G</b>

Alignments Select All Get selected sequences Distance tree of results

GENE ID: 1115793 zitR | zinc transport transcription regulator
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)

Score = 291 bits (744), Expect = 2e-80, Method: Compositional matrix adjust. Identities = 145/145 (100%), Positives = 145/145 (100%), Gaps = 0/145 (0%)

```
Query1MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLAEQISTNAKIAEKL<br/>MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLAEQISTNAKIAEKL<br
```

Query 121 LGNKFTDEEQEVISKFLSALTEEFQ 145 LGNKFTDEEQEVISKFLSALTEEFQ Sbjct 121 LGNKFTDEEQEVISKFLSALTEEFQ 145

GENE ID: 4433026 LACR 2420 | transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)

Score = 268 bits (685), Expect = 2e-73, Method: Compositional matrix adjust. Identities = 130/145 (89%), Positives = 141/145 (97%), Gaps = 0/145 (0%)

```
{\tt MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL\ MSLANQIDQFLG\ IMQFAENKHEILLG+CES+VKLTSTQEHILM+LA\ ++STNA+IAE+L\ }
Query
                                                                                       60
             MSLANQIDQFLGAIMQFAENKHEILLGECESNVKLTSTQEHILMILAAEVSTNARIAEQL
Sbjct 1
                                                                                       60
             KISPAAVTKALKKLOEOELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYOE
Query
       61
                                                                                       120
             KISPAAVTKALKKLOEOELIKSSRATNDERVVLWSLTEKAVPVAKEHA HHEKTLSTYOE
       61
             KISPAAVTKALKKLÕEÕELIKSSRATNDERVVLWSLTEKAVPVAKEHAAHHEKTLSTYÕE
                                                                                       120
Sbict
       121
             LGNKFTDEEQEVISKFLSALTEEFQ
Ouerv
                                             145
             LG+KFTDEEQ+VIS+FLS LTEEF+
             LGDKFTDEEQKVISQFLSVLTEEFR
Sbjct
```

GENE ID: 4799067 zitR | transcriptional regulator of the zit operon
[Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)

Score = 267 bits (682), Expect = 3e-73, Method: Compositional matrix adjust. Identities = 129/145 (88%), Positives = 141/145 (97%), Gaps = 0/145 (0%)

```
Query 1
             MSLANQIDQFLGTIMQFAENKHEILLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKL
             MSLANQIDQFLG IMQFAENKHEILLG+CES+VKLTSTQEHILM+LA ++STNA+IAE+L
Sbjct
             MSLANQIDQFLGAIMQFAENKHEILLGECESNVKLTSTQEHILMILAAEVSTNARIAEQL
                                                                                    60
       61
             KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQE
                                                                                    120
Query
             KISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA+PVAKEHA HHEKTLSTYQE
                                                                                    120
Sbjct
       61
             KISPAAVTKALKKLÕEÕELIKSSRATNDERVVLWSLTEKAIPVAKEHAAHHEKTLSTYQE
             LGNKFTDEEQEVISKFLSALTEEFQ 145
LG+KFTDEEQ+VIS+FLS LTEEF+
LGDKFTDEEQKVISQFLSVLTEEFR 145
Query
       121
Sbjct 121
```

>ref[YP 001033131.1]  $\fine G$  MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] Length=295

```
GENE ID: 4797387 rmaB | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 53.9 bits (128), Expect = 6e-09, Method: Compositional matrix adjust. Identities = 38/101 (37%), Positives = 54/101 (53%), Gaps = 3/101 (2%)
                 ILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAV 101
++ L E TNA+IAE L I P++VT +K+L+E E++ + ND+RV LTEK
LVELWNEDGLTNAEIAELLDIKPSSVTTQVKQLEEAEMVIRKQDENDKRVNRIFLTEKGR 106
Ouerv
Sbjct
          47
                 102
Query
                 EAQETROTMHNDISETI--FGN-LTDEEQEQLANLMEKLVE
Sbjct 107
>ref|YP_808725.1|  transcriptional regulator [Lactococcus lactis subsp. cremoris
SK11]
Length=292
GENE ID: 4432217 LACR 0742 | transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 51.2 bits (121), Expect = 4e-08, Method: Compositional matrix adjust. Identities = 35/91 (38%), Positives = 50/91 (54%), Gaps = 3/91 (3%)
                 TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHH 111
TNA+IAE L I P++VT +K+L+E E++ + ND+RV LTEK + T H
Query 52
                 TNAEIAELLDIKPSSVTAQVKQLEEAEMVIRKQDENDKRVNRIFLTEKGREAQETRDTMH
Sbjct
          57
                 Ouerv
         112
Sbjct 117
>ref|NP 266864.1|  transcription regulator [Lactococcus lactis subsp. lactis Il1403] Length=791
GENE ID: 1114333 rmaB | transcription regulator
[Lactococcus lactis subsp. lactis I11403] (10 or fewer PubMed links)
 Score = 51.2 bits (121), Expect = 4e-08, Method: Compositional matrix adjust. Identities = 35/91 (38%), Positives = 50/91 (54%), Gaps = 3/91 (3%)
                 TNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHH 111
TNA+IAE L I P++VT +K+L+E E++ + ND+RV LT+K + T H
TNAEIAELLDIKPSSVTAQVKQLEEAEMVIRKQDENDKRVSRIFLTDKGREAQETRDTMH 116
Query 52
Sbjct
          57
                 112
Sbjct 117
>ref|YP 808147.1| 🖸 transcriptional regulator [Lactococcus lactis subsp. cremoris
SK11]
Length=169
GENE ID: 4432499 LACR 0099 | transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 50.4 bits (119), Expect = 7e-08, Method: Compositional matrix adjust. Identities = 27/98 (27%), Positives = 55/98 (56%), Gaps = 0/98 (0%)
                 HILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA 100
HIL L ++ T ++A KL ++ VT+A++ L + + + + + + ND++ + + +T K
Query
          41
                  HILSALTKKDLTGIELATKLSVTRGGVTRAVQNLIKHQFLTTYQADNDKKKIYYHITTKG
          53
Sbict
                 VPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS
          101
Query
                 + VA H H+ ++K++E++I FLS
LKVASIHDKMHKIMDLKLGQIFDKYNENEKSIILNFLS
Sbict 113
GENE ID: 4798609 rmaD | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 49.3 bits (116), Expect = 1e-07, Method: Compositional matrix adjust. Identities = 26/98 (26%), Positives = 55/98 (56%), Gaps = 0/98 (0%)
                 HILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA 100
HIL L ++ T ++A KL ++ VT+A++ L + + + +A ND++ + + +T K
HILSALTKKDLTGIELATKLSVTRGGVTRAVQNLIKHQFLTTYQADNDKKKIYYHITTKG 115
Query
Sbjct 56
                 VPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS
                                                                            138
Query
         101
                 + VA H H+ ++ +K+ + ++ +I FLS
LKVASIHDKMHKIMDLKLGQIFDKYNENDKSIILNFLS
Sbict
          116
```

```
>ref[NP_266269.1|  transcription regulator [Lactococcus lactis subsp. lactis Il1403]
Length=T63
GENE ID: 1113719 rmaD | transcription regulator [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 46.2 bits (108), Expect = 1e-06, Method: Compositional matrix adjust. Identities = 26/98 (26%), Positives = 55/98 (56%), Gaps = 0/98 (0%)
                 HILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA 100
HIL L ++ T ++A KL ++ VT+A++ L + + + + + +D++ + + LT K
HILSALTKEDLTGIELATKLSVTRGGVTRAVQNLIKYQFLTTYQSESDKKKIFYHLTVKG 112
Sbjct 53
                 VPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLS
Query 101
                                             ++ +K+ ++E+ +I
                               H+
                RKVATIHDKMHKIMDIRLGQIFDKYNEQEKSIILSFLS
Sbict 113
>ref[NP_267628.1|  transcription regulator [Lactococcus lactis subsp. lactis Il1403]
Length=356
GENE ID: 1115129 rmaC | transcription regulator
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 41.2 bits (95), Expect = 4e-05, Method: Compositional matrix adjust. Identities = 29/100 (29%), Positives = 52/100 (52%), Gaps = 1/100 (1%)
                 LLGKCESDVKLTSTQEHILMLL-AEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSS 83
Ouerv 25
                 + K E + T+ ++M+L + + ++++ +LKI AAVT+ LK L+E+ L+K
MTAKFEKSTGYSITRYQLMMILKCKGRCSQSQLQNELKIDSAAVTRHLKLLEEKNLVKRE 79
Sbjct 20
                 RATNDERVVLWSLTEKAVPVAKEHATHHEKTLSTYQELGN
Ouery
         84
                                  +T+KA
                 RNKENNREVFVEITDKAKNDLALCAKKHDDSLDESQQILN
                                                                               119
Sbjct 80
>ref[YP 001032345.1] G MarR family transcriptional regulator [Lactococcus lactis subsp.
cremoris MG1363]
Length=139
GENE ID: 4798214 rmaC | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 38.9 bits (89), Expect = 2e-04, Method: Compositional matrix adjust. Identities = 25/92 (27%), Positives = 49/92 (53%), Gaps = 1/92 (1%)
                  LLGKCESDVKLTSTQEHILMLL-AEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSS 83
Query 25
                 + K E + T+ ++M+L + + ++ +LKI AAVT+ LK L+E+ L+K
MTAKFEKSTGFSITRYQLMMILKCKGRCSQTQLQNELKIDSAAVTRHLKLLEEKNLVKRQ 79
Sbjct
          20
                  RATNDERVVLWSLTEKAVPVAKEHATHHEKTL 115
Query 84
                  R ++ R V +T++A + A H+ ++
RNKDNNREVFVEITDEAKADLERCAREHDNSV
Sbjct 80
>ref[YP 808479.1] C transcriptional regulator [Lactococcus lactis subsp. cremoris
SK11]
Length=166
GENE ID: 4433047 LACR 0453 | transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 37.0 bits (84), Expect = 9e-04, Method: Compositional matrix adjust. Identities = 31/111 (27%), Positives = 54/111 (48%), Gaps = 13/111 (11%)
                  QEHILMLLAEQISTNAKIAEK----LKISPAAVTKALKKLQEQELIKSSRATNDERVVL 93
Q IL +L E N+K+ +K L + P + ++ +KKL++++ I + D+R +
QGQILNILME----NSKMTQKNLVAQLDMRPQSASEMIKKLEKKQFISRQKDAQDKRGFI 98
Ouerv
Sbjct
          43
                  WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF 144
Query
          94
                  SLTEK V +E A E + FT+EE+ ++ + L E ISLTEKGKAVLEESAEQTELVPG---IMTSFTEEEKIEFARLIGKLQSEL 145
Sbict 99
>ref|NP 266463.1|  MarR family transcriptional regulator [Lactococcus lactis subsp. lactis Il1403]
Length=159
GENE ID: 1113917 napB | MarR family transcriptional regulator [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 36.2 bits (82), Expect = 0.001, Method: Compositional matrix adjust. Identities = 23/84 (27%), Positives = 46/84 (54%), Gaps = 0/84 (0%)
                  KIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKT 114
Ouerv 55
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Annex 2
                K+A L ++ A +K KKL ++ L++S + +++ + LT++ + H + H+K
KLAHHLYMTRGAASKIAKKLLKKNLVESYQIPQNKKEIYFRLTKEGQEINDRHESLHQKF 119
                                                            +++ + + LT++
Sbjct
         60
               LSTYQELGNKFTDEEQEVISKFLS 138
Query 115
                Q + ++ TDE I KFL+
SEKDQVIFDELTDESVSNILKFLN 143
>ref[NP 266747.1| G transcription regulator [Lactococcus lactis subsp. lactis Il1403]
Length=T43
GENE ID: 1114211 rmaJ | transcription regulator [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 35.8 bits (81), Expect = 0.002, Method: Compositional matrix adjust. Identities = 23/65 (35%), Positives = 33/65 (50%), Gaps = 0/65 (0%)
                ILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAV 101
Query 42
                             T K+ E+L + ++ LK+L+ + I
                                                                         R+ DER V
                IAMLAIENKMTINKLGEELSLDSGTLSPLLKRLEAKGYIVRKRSDKDERSVELFLTDKGA
Sbjct
         40
         102
                PVAKE 106
Ouerv
                 V KE
                QVKKE 104
Sbjct 100
cremoris MG1363]
Length=157
GENE ID: 4798013 napB | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 35.4 bits (80), Expect = 0.002, Method: Compositional matrix adjust. Identities = 23/83 (27%), Positives = 45/83 (54%), Gaps = 0/83 (0%)
                KIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKT 114
Ouerv
                K+A L ++ A +K KKL ++ LI+S + +++ + + LT++ + H + H+K
KLAHHLYMTRGAASKIAKKLLKKNLIESYQIPQNKKEIYFRLTKEGQQINDRHESLHQKF 117
Sbjct
                LSTYQELGNKFTDEEQEVISKFL 137
Query 115
                      Q + ++ TDE
                SQNDQVIFDELTDEAVSNILEFL
Sbjct 118
>ref[NP 266898.1]  MarR family transcriptional regulator [Lactococcus lactis subsp. lactis T11403]
Length=158
GENE ID: 1114368 rmaA | MarR family transcriptional regulator [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 35.4 bits (80), Expect = 0.002, Method: Compositional matrix adjust. Identities = 22/66 (33%), Positives = 36/66 (54%), Gaps = 3/66 (4%)
                HILMLLAEQISTNA---KIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLT
+L L++ I N +I + L I ++V +A+K L E+E + R D+RV SLT
QLLCLISLYIKDNQSQEQITDDLSIDKSSVHRAIKGLIEKEYVSRVRDEKDKRVYRVSLT
         41
Query
Sbjct
         42
                EKAVPV 103
Query
         98
                 +KA
                QKARDI 107
Sbjct 102
>ref|YP 808372.1| G MarR family transcriptional regulator [Lactococcus lactis subsp.
cremoris SK11]
Length=157
GENE ID: 4434175 LACR 0343 | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 35.4 bits (80), Expect = 0.002, Method: Compositional matrix adjust. Identities = 23/83 (27%), Positives = 45/83 (54%), Gaps = 0/83 (0%)
                KIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEKT 114
K+A L ++ A +K KKL ++ LI+S + +++ + LT++ + H + H+K
KLAHHLYMTRGAASKIAKKLLKKNLIESYQIPQNKKEIYFRLTKEGQQINDRHESLHQKF 117
Query
Sbjct
         58
         115
                LSTYQELGNKFTDEEQEVISKFL
Query
                      Q + ++ TDE
                SONDOVIFDELTDEAVSNILEFL
>ref[YP_001031774.1]  transcriptional regulator [Lactococcus lactis subsp. cremoris
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MG1363] Length=166

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GENE ID: 4798362 llmg 0424 | transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 35.0 bits (79), Expect = 0.003, Method: Compositional matrix adjust. Identities = 29/111 (26%), Positives = 53/111 (47%), Gaps = 13/111 (11%)
                QEHILMLLAEQISTNAKIAEK-----LKISPAAVTKALKKLQEQELIKSSRATNDERVVL
Q IL +L E N+K+ +K L + P + ++ +KKL++++ I + D+R +
QGQILNILME----NSKMTQKNLVAQLDMRPQSASEMIKKLEKKQFISRQKDAQDKRGFI
Ouerv
Sbjct
         43
                WSLTEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTEEF
Query
         94
                SLTEK V +E + + FT+EE+ ++ + L E ISLTEKGKAVLEESTEQTGRVPG---IMTSFTEEEKIEFARLIGKLQSEL
Sbjct 99
Length=156
GENE ID: 4796958 llmg 0529 | putative transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 32.3 bits (72), Expect = 0.017, Method: Compositional matrix adjust. Identities = 18/62 (29%), Positives = 38/62 (61%), Gaps = 7/62 (11%)
Query 33 VKLTSTQE---HILMLLAE----QISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRA 85 +KL+S E ++L++LA + ++ +A +LK+SP+ + K +K L ++ L++S+
               MKLSSGWEQSVYVLLILARLPENRTMSSIALANRLKVSPSYLKKIIKSLVDEGLLRSTTG
Sbjct
               TN
Query
         86
                    87
Sbjct 61
               KN
>ref|YP 808573.1|  BadM/Rrf2 family transcriptional regulator [Lactococcus lactis subsp. Cremoris SK11]
Length=156
GENE ID: 4433945 LACR 0579 | BadM/Rrf2 family transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 32.3 bits (72), Expect = 0.018, Method: Compositional matrix adjust. Identities = 18/62 (29%), Positives = 38/62 (61%), Gaps = 7/62 (11%)
               VKLTSTQE---HILMLLAE----QISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRA 85
+KL+S E ++L++LA + ++ +A +LK+SP+ + K +K L ++ L++S+
MKLSSGWEQSVYVLLILARLPENRTMSSIALANRLKVSPSYLKKIIKSLVDEGLLRSTTG 60
Sbjct 1
Query 86
               TN
                     87
Sbjct 61
              KN
                     62
>ref[NP 266714.1] G hypothetical protein L153086 [Lactococcus lactis subsp. lactis
I11403]
Length=156
GENE ID: 1114177 yffB | hypothetical protein [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 32.3 bits (72), Expect = 0.018, Method: Compositional matrix adjust. Identities = 18/62 (29%), Positives = 38/62 (61%), Gaps = 7/62 (11%)
Query 33 VKLTSTQE---HILMLLAE----QISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRA 85
               +KL+S E ++L++LA + ++ +A +LK+SP+ + K +K L ++ L++S+
MKLSSGWEQSVYVLLILARLPENRTMSSIALANRLKVSPSYLKKIIKSLVDEGLLRSTPG 60
Sbict
               TN
          86
Query
Sbjct 61
               KN
cremoris MG1363]
Length=160
 GENE ID: 4798315 rmaA | MarR family transcriptional regulator
[Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 31.6 bits (70), Expect = 0.031, Method: Compositional matrix adjust. Identities = 29/107 (27%), Positives = 52/107 (48%), Gaps = 10/107 (9%)
                 HILMLLAEQISTNA---KIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLT
Ouerv 41
                 +L L++ I N +I + L I ++V +A++ L E+E + R +D+R SLT QLLCLISLYIKDNQSQEQITDDLSIDKSSVHRAIRSLIEKEYVVRVRDEHDKRAYRVSLT
Sbjct
          42
                 EKAVPVAK--EHATHHEKTLSTYQELGNKFTDEEQEVISKFLSALTE 142
Query
                            E T + L
                                                              +E+E+ K L+ +T+
                                                   L
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>ref[YP_808765.1] G MarR family transcriptional regulator [Lactococcus lactis subsp.
cremoris SK11]
Length=155
GENE ID: 4432345 LACR 0786 | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 31.6 bits (70), Expect = 0.034, Method: Compositional matrix adjust. Identities = 20/65 (30%), Positives = 36/65 (55%), Gaps = 3/65 (4%)
              Query 42
        43
Sbict
        99
               KAVPV 103
Ouerv
              KARAI 107
Sbict 103
>ref[YP 808795.1] G MarR family transcriptional regulator [Lactococcus lactis subsp.
ref|YP_001033061.1|  MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363]
cremoris SK11]
Length=146
GENE ID: 4433123 LACR 0820 | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 31.2 bits (69), Expect = 0.044, Method: Compositional matrix adjust. Identities = 19/57 (33%), Positives = 31/57 (54%), Gaps = 3/57 (5%)
               Sbict 57
>ref[NP 266926.1]  MarR family transcriptional regulator [Lactococcus lactis subsp. lactis T11403]
Length=146
GENE ID: 1114397 rmaG | MarR family transcriptional regulator [Lactococcus lactis subsp. lactis Ill403] (10 or fewer PubMed links)
 Score = 31.2 bits (69), Expect = 0.045, Method: Compositional matrix adjust. Identities = 19/57 (33%), Positives = 31/57 (54%), Gaps = 3/57 (5%)
               Query 56
Sbjct 57
Length=154
GENE ID: 4797429 rmaH | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 28.9 bits (63), Expect = 0.18, Method: Compositional matrix adjust. Identities = 24/85 (28%), Positives = 41/85 (48%), Gaps = 1/85 (1%)
               AKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEK 113
Ouery
               +KIA+ S A + L L+ + LI + D R +L ++T+K VA+E
SKIAKFTHTSTARIATILNNLESKNLITREISRTDRRKILVAITDKGRRVAEEIRVEACS 110
Sbjct
         51
               TLS-TYQELGNKFTDEEQEVISKFL
Query 114
               L+ ++E+G + T+ E FL
NLARVFKEMGEERTESFIENFKMFL
>ref|YP 001032520.1|  MarR family transcriptional regulator [Lactococcus lactis subsp.
cremoris MG1363)
Length=127
GENE ID: 4797914 rmax | MarR family transcriptional regulator [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 28.1 bits (61), Expect = 0.31, Method: Compositional matrix adjust. Identities = 24/86 (27%), Positives = 36/86 (41%), Gaps = 6/86 (6%)
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AKIAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHATHHEK 113 + IA + S A V L L+E+ +I + D R +L LT+K KE T SDIARYIGASTARVANILNNLEEKGMISREISREDRRKILVFLTDKGRKETKERRTR--- 85

Query 54 Sbjct 29

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Query 114
              TLSTYQELGNKFTDEEQEVISKFLSA 139
              T + N F +E +F+ A
---TITRISNVFEAMGEERTQQFIEA 108
Sbict 86
>ref|NP 267064.1| G transcription regulator [Lactococcus lactis subsp. lactis Il1403]
Length=T54
 GENE ID: 1114538 rmaH | transcription regulator
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 28.1 bits (61), Expect = 0.35, Method: Compositional matrix adjust. Identities = 23/85 (27%), Positives = 41/85 (48%), Gaps = 1/85 (1%)
              Query 54
Sbjct
        51
              TLS-TYQELGNKFTDEEQEVISKFL 137
Query
        114
              L+ ++E+G + T+ E FL
NLARVFEEMGEERTESFIENFKLFL
>ref[YP_001032533.1] G transcriptional regulator [Lactococcus lactis subsp. cremoris
MG13631
Length=217
 GENE ID: 4798916 llmg 1224 | transcriptional regulator
[Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 28.1 bits (61), Expect = 0.38, Method: Compositional matrix adjust. Identities = 14/33 (42%), Positives = 24/33 (72%), Gaps = 1/33 (3%)
             EQISTNAKIAEKLKISPAAVTKALKKLQEQELI
E +S NA IA+KL +S + T+ +K+L ++EL+
Sbjct 23 ESVSINA-IAQKLSVSSPSATEMIKRLAKKELV
>ref|YP 809312.1|  Mn-dependent transcriptional regulator [Lactococcus lactis subsp.
cremoris SK11]
Length=217
GENE ID: 4432558 LACR 1369 | Mn-dependent transcriptional regulator [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 28.1 bits (61), Expect = 0.38, Method: Compositional matrix adjust. Identities = 14/33 (42%), Positives = 24/33 (72%), Gaps = 1/33 (3%)
Query 48 EQISTNAKIAEKLKISPAAVTKALKKLQEQELI
             E +S NA IA+KL +S + T+ +K+L ++EL+
ESVSINA-IAQKLSVSSPSATEMIKRLAKKELV
        23
>ref|NP 267461.1| 🖸 transcription regulator [Lactococcus lactis subsp. lactis Il1403]
Length=T56
 GENE ID: 1114954 rmaF | transcription regulator
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 27.7 bits (60), Expect = 0.42, Method: Compositional matrix adjust. Identities = 15/45 (33%), Positives = 24/45 (53%), Gaps = 0/45 (0%)
               IAEKLKISPAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKA 100
Ouerv
              + E L IS A+ + L\overline{+}E\overline{+}EL+ ND+R+ LTE+ LLEVLDISKQALNGPMNDLKEKELVHFKPNENDKRIKQLYLTEQG
Sbjct 63
>ref|NP 267696.1| G transcription regulator [Lactococcus lactis subsp. lactis Il1403]
GENE ID: 1115197 rmaI | transcription regulator [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 27.3 bits (59), Expect = 0.61, Method: Compositional matrix adjust. Identities = 32/142 (22%), Positives = 60/142 (42%), Gaps = 18/142 (12%)
               Query 11
Sbjct
               PAAVTKALKKLQEQELIKSSRATNDERVVLWSLTEKAVPVAKEHAT---HHEKTLSTYQ 119
+ T+ L++++++LIK + D R LTE+ E T H++K L+
Query
        64
        68
               RSTTTEILORMEKROLIKRKAS PTDAROKS VELTEEGKOYLPEIRTY I QGHNQKALAG--
Sbjct
        120
              ELGNKFTDEEQEVISKFLSALT 141
Ouerv
                      + EE + KFL+
              ----LSAEEIAAVEKFLNNFS 142
Sbict 126
```

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>ref[NP_266753.1] C NADPH-flavin oxidoreductase [Lactococcus lactis subsp. lactis
1114031
Length=216
 GENE ID: 1114217 yfiJ | NADPH-flavin oxidoreductase
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
Score = 27.3 bits (59), Expect = 0.67, Method: Compositional matrix adjust. Identities = 11/29 (37%), Positives = 18/29 (62%), Gaps = 0/29 (0%)
              THHEKTLSTYQELGNKFTDEEQEVISKFL
                           QE+ N T
               +HH+K+
              SHHQKSTDWTQEMSNFLTKPRREDVAKFL
>ref[NP 267412.1] G metalloregulator [Lactococcus lactis subsp. lactis Il1403]
Length=217
GENE ID: 1114905 ymiA | metalloregulator [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 26.6 bits (57), Expect = 1.1, Method: Compositional matrix adjust. Identities = 13/31 (41%), Positives = 23/31 (74%), Gaps = 1/31 (3%)
Query 50 ISTNAKIAEKLKISPAAVTKALKKLQEQELI
              +S NA IA+KL +S + T+ +K+L ++EL+
Sbjct 25 VSINA-IAQKLSVSSPSATEMIKRLAKKELV
>ref|YP 001031697.1|  putative cobalt ABC transporter ATP-binding protein [Lactococcus lactis subsp. cremoris MG1363]
Length=565
GENE ID: 4798558 cbiO | putative cobalt ABC transporter ATP-binding protein
[Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 26.2 bits (56), Expect = 1.2, Method: Compositional matrix adjust. Identities = 12/27 (44%), Positives = 17/27 (62%), Gaps = 0/27 (0%)
               KIAEKLKISPAAVTKALKKLQEQELIK
Ouerv 55
                       ISP ++TKA
               ++A K
               QLARKADISPISLTKAFINFQNQERLK
Sbjct 537
GENE ID: 1114342 qor | quinone oxidoreductase [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 25.4 bits (54), Expect = 2.4, Method: Compositional matrix adjust. Identities = 10/19 (52%), Positives = 11/19 (57%), Gaps = 0/19 (0%)
               THHEKTLSTYQELGNKFTD
                             ELG KF
Sbjct 193 NHHEKLVPQVHELGFKFVD 211
>ref[NP 268402.1| G alkylphosphonate uptake protein [Lactococcus lactis subsp. lactis
I11403]
Length=114
GENE ID: 1115923 phnA | alkylphosphonate uptake protein [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 25.4 bits (54), Expect = 2.4, Method: Compositional matrix adjust. Identities = 14/41 (34%), Positives = 19/41 (46%), Gaps = 0/41 (0%)
               TEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFL 137
Query 97
               T K + + E+ T E GN+FT EE E KF+
TPKCIHCSSEYTYELSDTSFGCSECGNEFTLEEIEAAGKFI
Sbict 3
>ref|YP 808209.1|  transcriptional repressor CodY [Lactococcus lactis subsp. cremoris
SK11)
Length=262
 GENE ID: 4434653 LACR 0168 | transcriptional repressor CodY
[Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 24.6 bits (52), Expect = 3.4, Method: Compositional matrix adjust. Identities = 10/27 (37%), Positives = 19/27 (70%), Gaps = 0/27 (0%)
               IAEKLKISPAAVTKALKKLQEQELIKS 82
Query 56
               IA+K+I++AL+KL+
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>ref[YP 001031533.1]  transcriptional repressor CodY [Lactococcus lactis subsp. cremori
Length=262
 GENE ID: 4797198 cody | transcriptional repressor CodY
[Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 24.6 bits (52), Expect = 3.4, Method: Compositional matrix adjust. Identities = 10/27 (37%), Positives = 19/27 (70%), Gaps = 0/27 (0%)
Query 56
               IAEKLKISPAAVTKALKKLQEQELIKS 82
IA+K+ I+ + + AL+KL+ +I+S
Sbjct 210 IADKIGITRSVIVNALRKLESAGVIES 236
>ref[NP 266317.1]  transcriptional repressor CodY [Lactococcus lactis subsp. lactis
1114031
Length=262
GENE ID: 1113769 codY | transcriptional repressor CodY [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 24.6 bits (52), Expect = 3.5, Method: Compositional matrix adjust. Identities = 10/27 (37%), Positives = 19/27 (70%), Gaps = 0/27 (0%)
               IAEKLKISPAAVTKALKKLQEQELIKS
Ouerv 56
               IA+K+I++AL+KL+
Sbjct 210 IADKIGITRSVIVNALRKLESAGVIES
>ref[NP 266838.1]  intercellular adhesion protein [Lactococcus lactis subsp. lactis
1114031
Length=276
 GENE ID: 1114306 icaB | intercellular adhesion protein
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 24.6 bits (52), Expect = 3.6, Method: Compositional matrix adjust. Identities = 14/46 (30%), Positives = 21/46 (45%), Gaps = 0/46 (0%)
               LMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQELIKSSRATND
Query 43
L L Q K+ + + S + K +KK+ E L+ S TND Sbjct 126 LDLPFTQFIITGKVGQTIDGSQMSTWKEIKKMNENPLVTSGLHTND
>ref|NP 266710.1|  transcription regulator [Lactococcus lactis subsp. lactis Il1403] Length=247
 GENE ID: 1114173 yfeA | transcription regulator
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 24.6 bits (52), Expect = 3.6, Method: Compositional matrix adjust. Identities = 16/44 (36%), Positives = 25/44 (56%), Gaps = 5/44 (11%)
Query 35 LTSTQEHILMLLAEQIS----TNAKIAEKLKISPAAVTKALKK LTS + + I L E + T A+IAE +SP+++ + LKK Sbjct 4 LTSVEIYIWNYLEENKAKIIQMTVAQIAESAHVSPSSIIRTLKK
                                                                            73
>ref[NP 266438.1] G amino acid amidohydrolase [Lactococcus lactis subsp. lactis Il1403]
GENE ID: 1113892 yciA | amino acid amidohvdrolase [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 24.6 bits (52), Expect = 4.1, Method: Compositional matrix adjust. Identities = 11/41 (26%), Positives = 17/41 (41%), Gaps = 7/41 (17%)
                DQFLGT----IMQFAENKHEILLGKCESDVKLTSTQEH
D+F G + A N+H + G CE ++ T H
Sbjct 147 DEFYGLHVRPDLKVGDIATNQHTLFAGTCEVELSFIGTGGH 187
>ref|YP 808446.1| 🖸 lysyl-tRNA synthetase [Lactococcus lactis subsp. cremoris SK11]
Length=494
GENE ID: 4433352 lyss | lysyl-tRNA synthetase [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 24.6 bits (52), Expect = 4.3, Method: Composition-based stats. Identities = 12/34 (35%), Positives = 20/34 (58%), Gaps = 0/34 (0%)
                EKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQE 131
Query 98
```

E+A +AKEH H EK ++ + N+F ++ E

```
>ref[YP 001031741.1] 🖸 lysyl-tRNA synthetase [Lactococcus lactis subsp. cremoris MG1363]
 GENE ID: 4799123 lyss | lysyl-tRNA synthetase
[Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
Score = 24.3 bits (51), Expect = 4.7, Method: Composition-based stats. Identities = 12/34 (35%), Positives = 20/34 (58%), Gaps = 0/34 (0%)
                  EKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQE
Query
                 E+A +AKEH H EK ++ + N+F ++ E
EEATALAKEHDIHVEKHFTSVGHIINEFFEKYVE
Sbjct 340
>ref[NP 266529.1| G lysyl-tRNA synthetase [Lactococcus lactis subsp. lactis Il1403]
Length=494
GENE ID: 1113984 lyss | lysyl-tRNA synthetase [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 24.3 bits (51), Expect = 4.7, Method: Composition-based stats. Identities = 12/34 (35%), Positives = 20/34 (58%), Gaps = 0/34 (0%)
Query 98
                  EKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQE 131
E+A +AKEH H EK ++ + N+F ++ E
Sbjct 340 EEATALAKEHDIHVEKHFTSVGHIINEFFEKYVE
>ref|YP 001031907.1| G NADPH-flavin oxidoreductase [Lactococcus lactis subsp. cremoris
MG13631
Length=251
GENE ID: 4798593 llmg 0559 | NADPH-flavin oxidoreductase [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 24.3 bits (51), Expect = 5.2, Method: Compositional matrix adjust. Identities = 9/29 (31%), Positives = 19/29 (65%), Gaps = 0/29 (0%)
Query 109 THHEKTLSTYQELGNKFTDEEQEVISKFL 137
                  +HH+K+ + QE+ + T+ +E ++ FL
SHHQKSTNWSQEMSDFLTNPRREDLTDFL
Sbjct 215
>ref|YP 808603.1|    NADPH-flavin oxidoreductase [Lactococcus lactis subsp. cremoris
SK11]
Length=251
  GENE ID: 4433732 LACR_0613 | NADPH-flavin oxidoreductase
 [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 24.3 bits (51), Expect = 5.5, Method: Compositional matrix adjust. Identities = 9/29 (31%), Positives = 19/29 (65%), Gaps = 0/29 (0%)
Query 109 THHEKTLSTYQELGNKFTDEEQEVISKFL 137
+HH+K+ + QE+ + T+ +E ++ FL
Sbjct 215 SHHQKSTNWSQEMSDFLTNPRREDLADFL 243
>ref[NP_268293.1]  exported serine protease [Lactococcus lactis subsp. lactis Il1403]
Length=408
 GENE ID: 1115813 htrA | exported serine protease [Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
  Score = 23.9 bits (50), Expect = 6.1, Method: Compositional matrix adjust. Identities = 17/59 (28%), Positives = 34/59 (57%), Gaps = 5/59 (8%)
Query 15 MQFAENKHEI--LLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKAL
+ FA +++ ++ K E+D K++ I M+ Q+STN + +LK+ P++VT +
Sbjct 271 LGFAIPSNDVVNIINKLEADGKISRPALGIRMVDLSQLSTND--SSQLKL-PSSVTGGV
                                                                                                                71
 >ref|YP 001033660.1|  housekeeping protease [Lactococcus lactis subsp. cremoris MG1363]
 GENE ID: 4797497 htrA | housekeeping protease [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
  Score = 23.9 bits (50), Expect = 6.5, Method: Compositional matrix adjust. Identities = 17/59 (28%), Positives = 34/59 (57%), Gaps = 5/59 (8%)
                   \begin{array}{lllllllllllll} \texttt{MQFAENKHEI--LLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKAL} \\ + & \texttt{FA} & +++ & ++ & \texttt{K} & \texttt{E+D} & \texttt{K++} & \texttt{I} & \texttt{M+} & \texttt{Q+STN} & + & +\texttt{LK+} & \texttt{P++VT} & + \\ \end{array} 
                                                                                                                 71
 Query 15
                  + FA +++ ++ K E+D K++ I M+ Q+STN + +LK+ P++VT +
LGFAIPSNDVVNIINKLETDGKISRPALGIRMVDLSQLSTND--SSQLKL-PSSVTGGV 325
 Sbjct 270
```

```
>ref|YP 811995.1| G trypsin-like serine protease [Lactococcus lactis subsp. cremoris
SK11]
Length=407
GENE ID: 4432303 LACR 2439 | trypsin-like serine protease [Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 23.9 bits (50), Expect = 6.6, Method: Compositional matrix adjust. Identities = 17/59 (28%), Positives = 34/59 (57%), Gaps = 5/59 (8%)
              MQFAENKHEI--LLGKCESDVKLTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKAL
                       +++ ++ K E+D K++
                                                          Q+STN
                                                                   + +LK+ P++VT
                                                  I M+
             LGFAIPSNDVVNIINKLETDGKISRPALGIRMVDLSQLSTND--SSQLKL-PSSVTGGV 325
Sbict 270
>ref[NP 268332.1| G hypothetical protein L35545 [Lactococcus lactis subsp. lactis
I11403]
 ref|YP 812032.1|  hypothetical protein LACR_2483 [Lactococcus lactis subsp. cremoris
SK11]
 ref|YP 001033699.1| G hypothetical protein llmg_2459 [Lactococcus lactis subsp. cremori
MG1363]
Length=82
GENE ID: 1115852 ywfB | hypothetical protein
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 23.5 bits (49), Expect = 7.5, Method: Compositional matrix adjust. Identities = 13/29 (44%), Positives = 14/29 (48%), Gaps = 0/29 (0%)
              125
Query 97
              TEDGKKEAKEAAIRYESRLDAYOFLOGKF
Shict 34
>ref[NP 268337.1| G hypothetical protein L39650 [Lactococcus lactis subsp. lactis
I11403]
Length=926
GENE ID: 1115857 ywfG | hypothetical protein
[Lactococcus lactis subsp. lactis Il1403] (10 or fewer PubMed links)
 Score = 23.5 bits (49), Expect = 7.7, Method: Composition-based stats. Identities = 11/25 (44%), Positives = 13/25 (52%), Gaps = 0/25 (0%)
              YQELGNKFTDEEQEVISKFLSALTE 142
Query 118
             Y E G KF D+ Q I + LTE
YPEDGTKFADDPQHYIVRLKHGLTE 581
MG13631
Length=430
GENE ID: 4797767 comFA | superfamily II DNA/RNA helicase [Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 23.5 bits (49), Expect = 7.8, Method: Compositional matrix adjust. Identities = 17/46 (36%), Positives = 26/46 (56%), Gaps = 1/46 (2%)
              KLTSTQEHILMLLAEQISTNAKIAEKLKISPAAVTKALKKLQEQEL
Ouerv 34
KLT QE I L +QI+ N K+ + ++ A T+ + +L EQ L
Sbjct 105 KLTENQEKISNALCQQITNNQKLLVQ-AVTGAGKTEMIYQLIEQIL 149
>ref[YP 001033122.1] G quinone oxidoreductase [Lactococcus lactis subsp. cremoris MG1363
Length=\overline{3}28
 GENE ID: 4798878 qor | quinone oxidoreductase
[Lactococcus lactis subsp. cremoris MG1363] (10 or fewer PubMed links)
 Score = 23.5 bits (49), Expect = 9.3, Method: Compositional matrix adjust. Identities = 9/18 (50%), Positives = 10/18 (55%), Gaps = 0/18 (0%)
              HHEKTLSTYQELGNKFTD 127
                           ELG KF
Sbjct 194 HHENLVPQVHELGFKFVD 211
>ref|YP 808733.1| G quinone oxidoreductase [Lactococcus lactis subsp. cremoris SK11]
Length=\overline{3}28
 GENE ID: 4432226 LACR 0751 | quinone oxidoreductase
[Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
```

```
Annex 2
 Score = 23.5 bits (49), Expect = 9.3, Method: Compositional matrix adjust. Identities = 9/18 (50%), Positives = 10/18 (55%), Gaps = 0/18 (0%)
Query 110 HHEKTLSTYQELGNKFTD 127
HHE + ELG KF D
Sbjct 194 HHENLVPQVHELGFKFVD 211
SK11]
Length=330
 GENE ID: 4405852 LACR C57 | hypothetical protein
[Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 23.5 bits (49), Expect = 9.5, Method: Compositional matrix adjust. Identities = 11/33 (33%), Positives = 22/33 (66%), Gaps = 0/33 (0%)
Query 53 NAKIAEKLKISPAAVTKALKKLQEQELIKSSRA 85
N K ++L S + ++A K+Q Q+L+K+S++
Sbjct 189 NIKDTQELDFSSSNFSEAQLKVQNQDLVKNSKS 221
SK11]
Length=114
 GENE ID: 4432134 LACR 2554 | hypothetical protein
[Lactococcus lactis subsp. cremoris SK11] (10 or fewer PubMed links)
 Score = 23.1 bits (48), Expect = 10.0, Method: Compositional matrix adjust. Identities = 13/41 (31%), Positives = 18/41 (43%), Gaps = 0/41 (0%)
              TEKAVPVAKEHATHHEKTLSTYQELGNKFTDEEQEVISKFL 137
T K + + E+ E GN+FT EE E KF+
TPKCIHCSSEYTYELSDMSFGCSECGNEFTLEEIEAAGKFI 43
Query 97
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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/525,449	11/09/2005	Isabelle Poquet	1169-034	3827	
20529 THE NATH LA	7590 07/07/2009 AW GROUP		EXAMINER		
112 South Wes	st Street		MARVICH, MARIA		
Alexandria, VA	1 22314		ART UNIT	PAPER NUMBER	
			1633		
			MAIL DATE	DELIVERY MODE	
		•	07/07/2009	PAPER	

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

# Application No. Applicant(s) Notice of Non-Compliant POQUET ET AL 10/525,449 Examiner Art Unit Amendment (37 CFR 1.121) 1633 MARIA B. MARVICH -- The MAILING DATE of this communication appears on the cover sheet with the correspondence address -is considered non-compliant because it has failed to meet the requirements of The amendment document filed on \_\_\_\_ 37 CFR 1.121 or 1.4. In order for the amendment document to be compliant, correction of the following item(s) is required. THE FOLLOWING MARKED (X) ITEM(S) CAUSE THE AMENDMENT DOCUMENT TO BE NON-COMPLIANT: 1. Amendments to the specification: A. Amended paragraph(s) do not include markings. B. New paragraph(s) should not be underlined. C. Other 2. Abstract: ☐ A. Not presented on a separate sheet. 37 CFR 1.72. ☐ B. Other 3. Amendments to the drawings: A. The drawings are not properly identified in the top margin as "Replacement Sheet," "New Sheet," or "Annotated Sheet" as required by 37 CFR 1.121(d). ☐ B. The practice of submitting proposed drawing correction has been eliminated. Replacement drawings showing amended figures, without markings, in compliance with 37 CFR 1.84 are required. C. Other 4. Amendments to the claims: ☐ A. A complete listing of all of the claims is not present. B. The listing of claims does not include the text of all pending claims (including withdrawn claims) C. Each claim has not been provided with the proper status identifier, and as such, the individual status of each claim cannot be identified. Note: the status of every claim must be indicated after its claim number by using one of the following status identifiers: (Original), (Currently amended), (Canceled), (Previously presented), (New), (Not entered), (Withdrawn) and (Withdrawn-currently amended). D. The claims of this amendment paper have not been presented in ascending numerical order. ⊠ E. Other: See Continuation Sheet. 5. Other (e.g., the amendment is unsigned or not signed in accordance with 37 CFR 1.4): For further explanation of the amendment format required by 37 CFR 1.121, see MPEP § 714. TIME PERIODS FOR FILING A REPLY TO THIS NOTICE: 1. Applicant is given no new time period if the non-compliant amendment is an after-final amendment or an amendment filed after allowance. If applicant wishes to resubmit the non-compliant after-final amendment with corrections, the entire corrected amendment must be resubmitted. 2. Applicant is given one month, or thirty (30) days, whichever is longer, from the mail date of this notice to supply the correction, if the non-compliant amendment is one of the following: a preliminary amendment, a non-final amendment (including a submission for a request for continued examination (RCE) under 37 CFR 1.114), a supplemental amendment filed within a suspension period under 37 CFR 1.103(a) or (c), and an amendment filed in response to a Quayle action. If any of above boxes 1. to 4. are checked, the correction required is only the corrected section of the non-compliant amendment in compliance with 37 CFR 1.121. Extensions of time are available under 37 CFR 1.136(a) only if the non-compliant amendment is a non-final amendment or an amendment filed in response to a Quayle action. Failure to timely respond to this notice will result in: Abandonment of the application if the non-compliant amendment is a non-final amendment or an amendment filed in response to a Quayle action; or Non-entry of the amendment if the non-compliant amendment is a preliminary amendment or supplemental amendment. /Maria B Marvich/ Primary Examiner, Art Unit 1633

Continuation of 4(e) Other: Clam 13 indicates that it is previously presented, however, amendment has been made to delete the phrase "in which TATAAT represents the -10 box of said promoter" without markings.

As well, the objections to the claims and rejections under 35 USC 112, first paragraph made in the office action mailed 10/7/08 have not been addressed in the repsonse mailed 4/7/09. For example, the objection to claim 16 and 18 for reference to a previous claim using the article "an" as opposed to "the" has not been addressed. by amendment or argument. Similarly, the objection to claims i.e. 20 for recitation of at least one has not been addressed. Recommendation has been made to use the article "the" as opposed to "at least one". Claim 15 has been objected to as being a duplicate of claim 12, however, applicants have not addressed this objection. Finally, the rejection under 35 USC 112, first paragraph has not been addressed.

NATH & ASSOCIATES, PLLC DBA: THE NATH LAW GROUP

8108

7/2009 DATE : Apr

: 8108 CHE AMOUNT :\$1,110.00 ACCOUNT: GENERAL - 28

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Patent Office Fee Serial No. 10/525,449

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